

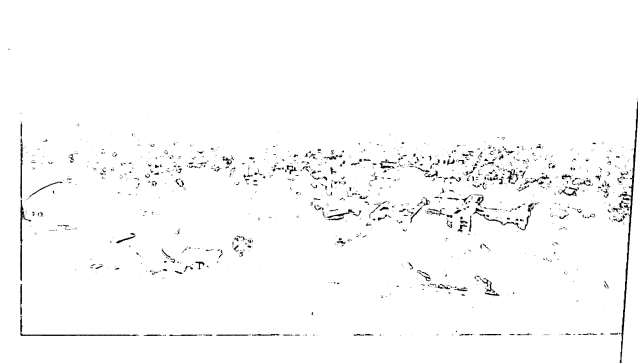
Lawrence Berkeley Laboratory University of California

CHEMICAL SCIENCES DIVISION

Reaction Dynamics and Photochemistry of Divalent Systems

H.F. Davis (Ph.D. Thesis)

May 1992



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REACTION DYNAMICS AND PHOTOCHEMISTRY OF DIVALENT SYSTEMS

H. Floyd Davis

Department of Chemistry University of California

and

Chemical Sciences Division Lawrence Berkeley Laboratory Berkeley, CA 94720 USA

May 1992

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REACTION DYNAMICS AND PHOTOCHEMISTRY OF DIVALENT SYSTEMS

H. Floyd Davis

Abstract

This thesis presents the results of molecular beams studies of bimolecular and unimolecular reactions. The rich chemistry seen in these systems is a direct consequence of the divalent nature of barium and oxygen atoms.

Chapter 1 discusses the reaction Ba + NO₂. Formation of the dominant BaO($^{1}\Sigma$) + NO products resulted primarily from decay of long-lived Ba $^{+}$ NO₂ collision complexes. Secondary mechanisms led to formation of forward scattered, internally excited BaO, and BaNO + O. Our results indicate that D_o(Ba-NO) = 65 \pm 20 kcal/mol.

The reactions of ground state and electronically excited Ba with water and alcohols are examined in Chapter 2. Reaction of $Ba(^1S) + H_2O$ led to $BaO + H_2$, whereas excited state $Ba(^1D) + H_2O$ reacted to form BaOH + H. Collisions between Ba and CH_3OH led to $BaOCH_3 + H$. The radical channels involve H-atom migration, and are promoted by electronic excitation of the incident Ba atom.

In Chapter 3, the reactions of Ba(1 S) with ClO₂ and O₃ are discussed. Again, direct and complex mechanisms were observed. Formation of BaCl + O₂ from concerted decomposition of long-lived Ba+ClO₂- collision complexes accounted for ~10% of the total reaction cross section. Although Ba + O₃ \rightarrow BaO + O₂ occurs primarily by direct reaction mechanisms, the secondary channel Ba + O₃ \rightarrow BaO₂ + O involved decay of long lived Ba+O₃- intermediates. We find that D_o(Ba-O₂) = 120 \pm 20 kcal/mol.

The photodissociation dynamics of NO₃ is explored in chapter 4. We confirm that visible excitation leads to formation of NO + O₂ and NO₂ + O. The wavelength dependence of the branching ratios is investigated. We find that $D_o(O-NO_2) = 48.55 \pm 0.16$ kcal/mole and calculate $\Delta H_f(NO_3) = 17.75 \pm 0.24$ kcal/mole (298K).

Chapter 5 discusses the photodissociation of OCIO in a molecular beam. Although $ClO(^2\Pi) + O(^3P)$ is dominant, we also observed formation of $Cl(^2P) + O_2$. The yield reaches a maximum of $3.9 \pm 0.8\%$ near 404nm. Mode specificity was seen in these branching ratios. The $Cl + O_2$ results from concerted unimolecular decomposition with a large release of product translational energy.

CHAPTER 1

Reactions of Ground State and Electronically Excited Ba with NO₂

ABSTRACT

Angular and velocity distributions of the neutral products resulting from the reaction Ba + NO₂ were measured using the crossed molecular beams method. Despite a large reaction exoergicity ($\Delta H = -61 \text{ kcal/mole}$), formation of the dominant ground state $BaO(^{1}\Sigma)$ + NO products results primarily from decay of long-lived $Ba^{+}NO_{2}^{-}$ collision complexes, even at incident collision energies as high as 59 kcal/mole or with electronic excitation of the Ba atom. A large fraction of the reaction exoergicity is channeled into product translational energy. This rather unusual behavior results from a large exit potential energy barrier for decay of the initially formed singly ionic Ba+NO₂intermediate to ground state doubly ionic Ba²⁺O²⁻. A secondary source of forward scattered, internally excited BaO results from a direct reaction without the involvement of long-lived intermediates. An additional minor channel, formation of BaNO + O is observed from ground state Ba + NO₂ at high collision energies by a direct reaction mechanism. Unlike the dominant BaO + NO channel which involves harpooning at the first ionic-covalent curve crossing, formation of BaNO from reaction of ground state Ba likely results from the small range of collision geometries which are able to avoid long range electron transfer. The BaNO + O channel was enhanced substantially by electronic excitation of the incident barium atom. However, BaNO from reactions of electronically excited Ba primarily resulted from decay of collision complexes, rather than from a direct mechanism.

I. INTRODUCTION

The early studies by Michael Polanyi and coworkers^{1,2} on the reactions of alkali metal atoms with halogen containing molecules in diffusion flames initiated a field of research that has led to profound insight into the reactions of monovalent species. The early observation that reaction rates were larger than those expected by gas kinetic theory led to the now famous "harpoon mechanism" 1-6 used to describe the general class of reactions initiated by long range electron transfer.⁴ In the 60's the advancement of crossed molecular beams techniques confirmed these early ideas and led to a detailed understanding of the reaction dynamics of monovalent atoms.^{5,6} In the reaction K + Br₂ → KBr + Br, the facile dissociation of newly formed diatomic halogen anions⁷ under the influence of alkali cations led to the "spectator stripping" mechanism with the product salt molecule forward scattered with respect to the incoming alkali atom. On the other hand, the pronounced backscattering of the KI product from the reaction K + CH₃I⁸ resulted from the need for approach of the alkali atom towards the halogen end of the molecule, followed by strong repulsive forces in the exit channel. These two prototype reactions illustrated two extremes in chemical dynamics.

Although most early studies were restricted to alkali atoms due to the ease of product detection using surface ionization,⁹ the development of the "universal" crossed molecular beams apparatus¹⁰ and optical techniques such as laser induced fluorescence¹¹

and chemiluminescence detection¹² permitted studies of a variety of other neutral atom-molecule reactions. More than two decades ago, BaO chemiluminescence was first observed from single collisions between Ba atoms and oxygen- containing triatomic molecules. ¹³⁻¹⁵ Due to the potential applications of these systems in the development of electronic transition chemical lasers, these and many subsequent studies focussed on the identification and relative yields of nascent chemiluminescent species. Although hopes for practical laser applications soon faded, fascination with the reactions continued primarily because the origin of the BaO emission spectrum was not well understood. ^{16,17} In most cases, the chemiluminescence yield was found to be pressure dependent: increasing pressure led to increased chemiluminescence. More than 50 papers ¹⁶ were devoted to these studies but most focussed on Ba + N₂O since the effect in that case was particularly dramatic—the chemiluminescence yield increased from 2.5% to 20% upon increasing the pressure from 1x10⁻⁴ Torr to 10 Torr.

There has been considerably less work devoted to the reaction Ba + NO₂, primarily because of the small photon yield. It was found that more than 98% of all BaO was formed in non-chemiluminescent states. ^{16,17} Careful measurements by several groups showed that the collision free yield of chemiluminescent BaO(A¹ Σ ⁺) molecules was only 0.2%, with the value reaching a maximum of \leq 2% at high pressure. According to Hsu and Pruett¹⁷, the very low high pressure yield could be attributed to the fact that the nascent BaO from Ba + NO₂ are almost all formed in the X¹ Σ ground state with only a small amount of vibrational energy. This dominant ground state BaO possessed insufficient internal energy for electronic excitation by secondary collisions.

Angular distributions of the BaO resulting from Ba + NO₂ were measured by Haberman, et al. 18 These experiments were rather primitive by present standards, employing effusive beams with broad beam velocity distributions, and no product velocity distributions could be obtained. They concluded that the BaO product was forward scattered with respect to the Ba beam with only a small fraction of the available energy appearing in BaO-NO translational recoil. Herm and coworkers, ¹⁹ using a similar apparatus studied the reactions of Sr and Ca (but not Ba) with NO2. Their results on the lighter alkaline earth species were consistent with the results of Haberman on Ba + NO₂. Based on the work from these two laboratories, it was concluded that the reactions of alkaline earth atoms with NO2 were analogous to the well-studied alkali-halogen systems.^{7,20} Due to the low ionization potential of Ba (5.2eV)²¹ and large electron affinity of NO₂ (2.3eV)²², the reaction is initiated by long range electron transfer, 4,13-15 consistent with a very large experimentally determined reaction cross section (>150Å²).¹³ By analogy with the alkali-halogen systems, it was argued that after long range electron transfer, the strong field of the approaching Ba⁺ led to immediate dissociation of NO₂⁻. The incoming Ba+ "picked up" the O ion with the newly formed BaO molecule continuing in nearly the same center-of-mass (CM) direction as the incident Ba atom. 18,19

This direct reaction mechanism contrasted the general behavior seen earlier by Ham and Kinsey²³ in scattering of alkali atoms such as Cs with SO₂, CO₂ and NO₂. Angular distributions measured using a surface ionization detector provided good evidence that collisions between alkali atoms and these triatomic oxidants led to formation of strongly bound ionic collision complexes (e.g. Cs+SO₂). The center-of-mass

angular distributions of the nonreactively scattered Cs showed forward-backward symmetry, indicating that the lifetime of the ionic intermediate exceeded several rotational periods.²⁴ Decay of a long-lived prolate collision complex is expected to result in a center-of-mass product angular distribution peaking at $\theta_{CM} = 0^{\circ}$ and 180°; this often leads to a characteristic laboratory angular distribution exhibiting two peaks. In the case of SO₂ and CO₂, chemical reaction was too endoergic to be possible; the observed Cs signal could only result from decay of complexes back to reactants. Although the reaction Cs + NO₂ → CsO + NO was thought to be slightly exoergic, it was not clear from that work whether or not a reaction actually occurred. Subsequently, Herm and Herschbach²⁵ observed formation of CsO with a large cross section from Cs + NO₂. The bimodal character of the observed CsO laboratory angular distribution was again strongly suggestive of the participation of long-lived Cs⁺NO₂⁻ reaction intermediates. However, they resisted drawing any definite conclusions because uncertainties remained in the analysis of the data for the reactive channel since the product translational energy distributions were not recorded. We have recently measured angular and velocity distributions for the reaction Na + $NO_2 \rightarrow NaO$ + NO using supersonic beams in our apparatus.²⁶ We observe strong NaO signal from decay of long-lived Na⁺NO₂⁻ intermediates. Based on this and the past nonreactive and reactive scattering results, there is little doubt that reaction of the heavier alkalis including Cs + NO₂ involve longlived reaction intermediates.²⁷

That alkalis react with NO₂ by formation of long-lived complexes is not surprising since the reactions are nearly thermoneutral, NO₂ is a stable ion, and the ion pair

 $Cs^+NO_2^-$ has a deep potential well. The conclusion that alkaline earth atoms react by a direct stripping mechanism, however, also seems very reasonable. The Ba⁺NO₂⁻ and $Cs^+NO_2^-$ singly charged ionic intermediates are expected to be similar; both should be bound by about 65 kcal/mole with respect to reactants.^{25,28} However, the reaction exoergicities are widely different: Ba+ $NO_2 \rightarrow BaO + NO$ is highly exoergic- $\Delta H = -61$ kcal/mole²⁹ (Table I), whereas $Cs + NO_2 \rightarrow CsO + NO$ is now known to be slightly endoergic ($\Delta H = +2$ kcal/mole).²⁹ The large difference is due to the divalent nature of barium: the ground state BaO molecule, best described as $Ba^2 + O^{2-30}$ is bound by 133 kcal/mole, whereas the Cs^+O^- binding energy is about half this value (~ 70 kcal/mole).²⁹ Based on thermodynamics alone, the large exoergicity of the Ba reaction is expected to lead to subpicosecond lifetimes for $Ba^+NO_2^{-31}$ with the BaO forward scattered; whereas the slight endoergicity of the reaction of cesium allows the $Cs^+NO_2^-$ reaction intermediate to survive many rotational periods before decaying to products or back to reactants.

Although velocity distributions were not measured in the previous alkaline earth + NO₂ crossed beams experiments, based on the product angular distributions it was concluded that the nascent products were formed with relatively low translational energy. The more recent laser induced fluorescence work indicated that the dominant ground state BaO was vibrationally cold. The following question arises: Where has the 61 kcal/mole reaction exoergicity been disposed? For a reaction apparently involving long range electron transfer with immediate dissociation of NO₂, there is no obvious reason why the reaction exoergicity should be exclusively channeled into vibrational excitation of the NO product. Although it is known that the products are

rotationally hot, it seems impossible to believe that so much energy could be preferentially funneled into rotation without the accompanying release of some of the excess energy into translation.

The goal of the present work is to probe the dynamics of the reactions of a divalent metal atom with oxygen containing molecules under single collision conditions. Barium was deemed to be the most suitable alkaline earth atom for several reasons. Owing to its relatively low second ionization potential (10ev)²¹ it is most likely to exhibit dynamical effects resulting from its divalency. Also, the first optically allowed electronic transition from the ground state (¹P ←¹S) is at a wavelength convenient to Ar+ laser pumped ring dye lasers whereas transitions for the lighter alkaline earths are at less accessible, shorter wavelengths.³² Here we report^{33a} on the reaction Ba + NO₂, and a later chapter will focus on Ba + ClO₂ and O₃.33b Although all three triatomic molecules are closely related, the behavior of each provides considerable insight into the reaction dynamics of divalent systems. Unlike all experiments undertaken to date, both the angular and velocity distributions of the products have been measured. Both beams are much better defined than in the past work from other laboratories, having a narrow velocity distribution characteristic of a supersonic expansion. Our results indicate that previous conclusions, that the Ba + NO₂ reaction is analogous to alkali atoms + halogens is not quite correct. We have found that the divalent nature of barium leads to rather interesting and unexpected reaction dynamics.

The primary advantage of the mass spectrometric detection method over chemiluminescence is that all chemical products, including those in the ground electronic

states are detected simultaneously. In addition to BaO + NO, a number of other chemical channels are thermodynamically possible from $Ba + NO_2$ (Table I). Mass detection provides the opportunity to observe such products that cannot easily be detected by optical techniques either because of experimental complexities or because their electronic spectrum is unknown. We illustrate this by the observation of BaNO in the present experiment. As far as we know, this is the first direct observation of a neutral metal mononitroxide in the gas phase.

II. EXPERIMENTAL

The universal crossed molecular beams apparatus is shown in Figure 1. A continuous, seeded supersonic beam of barium atoms was formed in a molybdenum oven source with separately heated barium reservoir and nozzle. Details of this source will be published elsewhere. The barium reservoir was heated to ~1050 °C, corresponding to a barium pressure of ~10 Torr, and pressurized with 300-700 Torr of an inert gas. The mixture was expanded through a 0.008" diameter nozzle heated to 1350 °C to minimize the formation of clusters. The beam was skimmed by a heated (1000°C) molybdenum skimmer located in the differential pumping region and then collimated to 2° full width by a set of defining slits.

The electronically excited barium atomic beam was produced by optically pumping the Ba(${}^{1}P \leftarrow {}^{1}S$) resonance either at or upstream of the interaction region using a Coherent 699-21 ring dye laser operating single frequency at 553nm with Rhodamine 560 dye. The fluorescence intensity was continuously monitored during the experiment

by imaging the light from the pumped region onto a photomultiplier tube using a commercial telephoto lens. Once the oven was stabilized, the fluorescence intensity remained constant to within a few percent over a period of several hours.

The metastable Ba(1.3D) population resulted from radiative decay from the 1P state $(\tau \sim 8.4 \text{ns})$. By pumping a 1cm length of the beam upstream of the collision zone at 100mW, it was found that 60% of the ground state ¹³⁸Ba population was depleted. This measurement was accomplished by splitting off a small portion (10mW) of the laser beam, passing it through an acoustoptic modulator and using the resulting beam to probe the ground state Ba population downstream of the primary pumping region. The 60% fluorescence depletion represents the fractional decrease in ground state population in the beam due to optical pumping to the long lived (~1sec) metastable (1,3D) states. This fraction was relatively insensitive to laser power, indicating that the transition was strongly saturated, as was expected at powers above 50mW. Our experiment, together with published34 Ba(1P) branching ratios indicates that by pumping upstream of the crossing region the ¹³⁸Ba in the beam consisted of 45% Ba(¹D), 15% Ba(³D) and 40% Ba(1S), with all other isotopes in the ground (1S) state. The reaction of Ba(1P) was studied by focussing the laser to 3mm diameter and pumping at the interaction region. Under these conditions, the ¹³⁸Ba consisted of 30% Ba(^{1,3}D), 35%Ba(¹P) and 35%Ba(¹S). Although the Ba(1P) atoms are well aligned, we have found no orbital alignment dependence for the neutral product channels from Ba + NO₂.35

The NO₂ beam was formed by passing a noble gas through liquid or solid NO₂ cooled in a thermostatted temperature bath. The seeded molecular beams were typically

5-20% NO_2 in He; the lowest collision energy was obtained using neat NO_2 vapor. In each case, the NO_2 beam nozzle was heated to 200-250°C to minimize the formation of N_2O_4 and larger clusters. Mass scans with the detector looking directly into the molecular beam indicated that the concentration of such species were negligible. To be certain, parallel experiments were run using 100° C and 250° C nozzle temperatures in He carrier gas. The same beam velocity was attained in each experiment by addition of Ne to the He in the higher temperature experiment. The BaO product angular distributions were found to be the same, indicating negligible contribution from reaction of N_2O_4 . If there was substantial contribution from reaction of N_2O_4 , its higher concentration in the lower temperature experiment should have led to an increase of product at larger scattering angles since the center-of-mass angle associated with Ba + N_2O_4 scattering is substantially greater than that for Ba + N_2O_4

The Ba and NO_2 beam velocity distributions were measured using the time-of-flight method. A 17.8cm diameter stainless steel wheel with four 1mm slots cut equally spaced around its circumference was mounted to the front of the detector. The wheel was spun at 300Hz and the modulated beam was sampled directly through a 0.003" aperture which could be slid into the entrance of the detector. The distance between the wheel and the effective center of the ionizer was determined experimentally to be 17. cm by expanding various inert carrier gasses through a room temperature nozzle. The beam velocity distributions were obtained by fitting the time-of-flight profiles using KELVIN^{37a}, a program which convolutes over the known apparatus functions in determining the beam speed (v) and speed ratio (v/ Δ v). Typical beam parameters are

shown in Table II. It should be noted that a very wide range of Ba beam velocities can be obtained by choosing different inert seed gasses; in all cases the velocity distributions are narrow, characteristic of a supersonic expansion. For reactive scattering studies, the 4-slot wheel was replaced with a 17.8cm diameter cross correlation wheel³⁸ photoetched by PCM products. The wheel has two 255 channel pseudorandom sequences of open and closed slots providing 50% transmission and a nominal 5μ s time resolution when spun at 392Hz.

During reactive scattering experiments, the beam velocities were periodically checked using the cross correlation wheel by moving the detector into the beam, as in the velocity measurements described above. We were especially careful to reproduce the Ba oven heating voltages and currents since small changes in Ba vapor pressure can have substantial effects on the average mass of the beam and hence its velocity when light carrier gasses are used.

Angular distributions for Ba and BaO were measured in a separate series of experiments with the time of flight wheel removed. The NO₂ beam was modulated at 150Hz using a tuning fork chopper—the (beam on) - (beam off) intensities were recorded for each detector angle in separate channels and subtracted to give the reactive scattering signal. In studies of the effects of electronic excitation, the laser beam was chopped at 3Hz in synchronization with the NO₂ beam modulation and four channels were recorded:

1) Laser on, NO₂ on; 2) Laser on, NO₂ off; 3) Laser off, NO₂ on; 4) Laser off, NO₂ off. Further details of this procedure may be found elsewhere. The BaNO angular distributions were obtained by integrating the area under the peak in the time of flight

spectra obtained at appropriate detector angles.

For each species (Ba, BaO or BaNO), the data was recorded with the mass spectrometer tuned to the parent ion mass. The dominant (71%) ¹³⁸Ba isotope-containing species was monitored in each case. Product velocity distributions were obtained by running parallel experiments at various detector angles with the laser on and off. In all cases, tuning the laser well away from the barium atomic resonance had the same effect as blocking the laser beam.

III. RESULTS AND ANALYSIS

A. $Ba(^1S) + NO_2 \rightarrow BaO + NO$

Product angular distributions and time-of-flight data were recorded with the mass spectrometer set at Ba⁺ (m/e=138) and BaO⁺ (m/e=154) for the above reaction at 4 collision energies in the range 5-59 kcal/mole. The angular distributions of BaO exhibited two distinct peaks; at all collision energies the peak near 0° was larger than the wide angle peak (Fig. 2). The observed signal was very strong; typically only 10 mins. of averaging was required at each angle for each time-of-flight spectrum. The Ba⁺ data at all angles wider than 10° was very similar to the BaO data, indicating that the signal obtained for Ba⁺ primarily results from fragmentation of BaO in the ionizer with a small contribution from nonreactive scattering. Our observations are consistent with a large reaction cross section and the previous conclusions of "reactions at every collision." ¹³

In the analysis of the BaO data, an important concern is the internal energy dependence of the fragmentation pattern of BaO upon 200V electron impact ionization.

This turned out to be an important factor for NaO in our recent study³⁹ of the reaction $Na + O_3 \rightarrow NaO + O_2$. In that case, the fragmentation ratio Na^+/NaO^+ increased substantially with internal excitation of NaO, and we included the dependence in the analysis. We recorded the Ba⁺ and BaO⁺ time-of-flight data from Ba + NO₂ at the center-of-mass angle, where products with the full range of internal energies contribute to the TOF spectrum. The shape of both spectra were identical, indicating that internal energy of the precursor BaO is not high enough in the present case to noticeably affect the fragmentation pattern. Similar conclusions have been made by others in the past. 19,40 It is important to note that our detector is not state specific; we have no way to unambiguously distinguish between different nascent electronic states of BaO unless their times of arrival are well separated. However, excited states are known to be a minor contribution to the total reaction cross section. They will radiate³⁰ during their flight to the ionizer $(T > 100\mu\text{sec})$, and will be detected with the same efficiency as nascent ground state products. Formation of electronically excited $NO(A^2\Sigma^+)$ is not thermodynamically possible at the collision energies employed in this experiment.²⁹

We now consider possible explanations for the observed product angular distribution. Most commonly, the characteristic forward-backward peaking observed here results from decay of long-lived collision intermediates. As was discussed by Miller, et. al., ²⁴ decay of a prolate complex with a lifetime greater than a few rotational periods (>1-2psec) is expected to lead to a forward-backward symmetric center-of-mass product angular distribution, $T(\theta)$. The degree of peaking at the poles (θ_{CM} =0° and 180°) with respect to θ_{CM} =90° provides information about the disposal of the total angular

momentum into product rotation. Upon examining the laboratory angular distributions, one might anticipate that the CM angular distribution was completely forward-backward symmetric with the measured angular distribution distorted in favor of the forward peak due to the transformation Jacobian relating the product intensities in the laboratory to that in the center-of-mass coordinate system:²⁴

$$I_L(\otimes,v)=I_{CM}(\theta,u)\frac{v^2}{u^2} \qquad (1)$$

For decay of long-lived complexes, the recoil velocity in the center-of-mass frame of reference (u) is independent of angle. However, as can be seen from the Newton diagram (Fig. 2), the laboratory center-of-mass angle is less than 45° in this experiment. As a consequence, at small lab angles the lab velocity (v) associated with a given center-of-mass velocity (u) is greatest in the forward direction, leading to enhancement of $I_L(\Theta,v)$.

The product angular distributions and time-of-flight spectra were fit using a forward convolution program^{37d} that assumes a separable form for the center-of-mass product flux distribution:

$$I_{CM}(\theta,E) = T(\theta) \times P(E)$$
 (2)

where $T(\theta)$ is the CM product angular distribution, and P(E) is the CM relative product translational energy distribution. Using a trial $T(\theta)$ and P(E), the program generates the lab frame angular distribution and time-of-flight spectra for each lab angle after

convoluting over the measured beam velocity distributions and known apparatus functions. The program compares these calculated values with the experimental data and utilizes the simplex method^{37f} to optimize the $T(\theta)$ and P(E) iteratively to find a global minimum in the difference between the calculated and experimental angular distribution and times-of-flights.

We initially attempted to fit the time-of-flight and angular distributions with a single uncoupled CM angular ($T(\Theta)$) and translational energy (P(E)) distribution. This approach is expected to be correct for a single channel involving decay of complexes with lifetimes of several rotational and many vibrational periods. An uncoupled $T(\Theta)$ and P(E) is also likely to be a reasonable approximation if 'osculating'⁴¹⁻⁴⁴ complexes were involved with lifetimes on the order of one rotational period. Although the assumption of separability of the angular and energy distribution allowed us to reproduce the qualitative features of the product time-of-flight and angular distributions, it was not possible to obtain a fit commensurate with the quality of the raw data, which was reproducible and of good signal-to-noise ratio (Fig. 3).

Our data indicates that the average translational energy release in the forward direction is smaller than in the backward hemisphere. This leads us to conclude that the assumption of a single uncoupled $T(\theta)$ and P(E) (i.e. a single mechanism involving complexes) is unsatisfactory. The existence of osculating complexes with relatively short lifetimes can be ruled out since it is known that product angular distributions from their decay will be rather sensitive to collision energy. If the complexes lived for ~ 1 rotational period at low collision energy, upon increasing the collision energy we should

observe substantially increased forward scattering as their statistical lifetimes decrease. Such a transition has been observed in other systems. 41-44 In the present case, however, we see no evidence for this effect at a collision energy of 31 kcal/mole. At a collision energy of 59 kcal/mole, although we do see an increased contribution from forward scattering, the best fit of the angular and time-of-flight distributions indicated that most BaO resulted from decay of long-lived collision complexes.

The data can only be consistent with two alternative and somewhat more complicated models. The first possibility, which we will subsequently show to be correct, is that two different reaction mechanisms are involved; the dominant channel involves long- lived complexes with large translational energy release, and a second (minor) forward scattered channel with less translational energy imparted to the products. Alternatively, it could be possible that two direct reaction mechanisms are involved; one leading to forward scattering with low product translation and a second leading to backscattering with larger recoil energy. Although a direct reaction with forward scattered BaO is not at all surprising, the second model also requires direct backscattered BaO with large translational energy release. Although direct backscattered products are known to result from some bimolecular reactions, most notably Ba + N₂O → BaO + N_2^{45a} , or Ba + CH₃I \rightarrow BaI + CH₃^{45b}, the behavior in those cases is attributable to steric effects. Due to the necessity for attack on sterically hindered atoms such as O in linear N₂O or I in CH₃I, successful reaction requires approach of the metal atom from a direction near the symmetry axes (C_{∞v} or C_{3v} for N₂O or CH₃I, respectively). A direct reaction followed by strong repulsion in the exit channel leads to a "rebound" mechanism

with the metal atom reversing direction in the CM frame after formation of the chemical bond. Since NO₂ is a bent molecule, there appears to be very few collision geometries which could lead to such backscattering; such geometries should be far too rare to be a major source of backscattered BaO.

The reaction Ba + $NO_2 \rightarrow BaO_2 + N$ is thermodynamically possible: based on our measure of $D_o(Ba-O_2)^{33a,b}$, the reaction is calculated to be exoergic by 12 kcal/mole. Such a reaction (if it occurred at all) would most likely result from attack by the Ba atom near the C_{2v} axis between the two oxygen atoms, followed by concerted formation of BaO_2 . Since this might be expected to lead to backscattered BaO_2 , a possible source of strong signal could be from fragmentation of the peroxide to BaO^+ upon electron impact ionization. We discount this possibility in the present case since the maximum recoil velocity for BaO_2 is restricted to be very low due to the small exoergicity for the BaO_2 + N reaction and the light N recoil partner. The backscattered BaO that we observe experimentally has a much larger translational energy than would be possible from the $BaO_2 + N$ reaction. Moreover, we have characterized BaO_2 — it yields a parent ion peak from a more exoergic reaction ($Ba + O_3$). 33a,b Yet we cannot detect any BaO_2^+ from Ba + NO_2 .

Ruling out a direct reaction leading to backscattered BaO, we now return to the alternative explanation, that a direct forward scattered product is superimposed on a second channel involving long-lived complexes. We have analyzed the data based on this model. As shown in Fig. 4, two independent reaction channels were used in the fits; the first channel (channel 1) was constrained to forward-backward symmetry. This was the

dominant channel at all collision energies. The second channel (channel 2) is a relatively minor component and is forward scattered. The best fit of the data obtained at a mean collision energy of 12.5 kcal/mole is shown in Figs. 2 and 3. The first channel, corresponding to the decay of long-lived complexes, leads to larger translational energy release than the second channel. Upon increasing the incident reactant collision energy, the product angular and time-of-flight spectra become more compressed, with somewhat greater uncertainties in the translational energy distributions. However, the backward peak in the product angular distribution can be clearly seen at $E_{coll} = 31$ kcal/mole (Fig. 5), and the best fit of the angular distribution and TOF data (Fig. 6) even at the highest collision energy (59 kcal/mole) requires that the dominant channel exhibit forwardbackward symmetry in the CM product angular distribution (Fig. 7). The relative contributions from the two channels remained essentially constant to within experimental uncertainty over the 5-31 kcal/mole collision energy range. The BaO product flux contour maps based on the optimized P(E) and T(θ) at 12.5 kcal/mole are shown in Figs. 8,9. The forward scattered BaO component (channel 2) accounted for $20\pm5\%$ of the total BaO cross section at collision energies in the range 5-31 kcal/mole. The forward scattered component increased at 59 kcal/mole to 40±10%.

B. $Ba(^{1,3}D, ^{1}P) + NO_{2} \rightarrow BaO + NO$

The BaO angular distributions shown in Fig. 10 show the effect of electronic excitation of the Ba atoms at the interaction region. The beam contained 30% Ba (1,3D), 35% (1P) and 35% (1S). Although the effect was rather small, the BaO intensity was

found to increase at lab angles smaller than 25° and greater than 50°. The signal decreased slightly between 25° and 50°. The Newton diagram (Fig. 10) shows circles for the maximum BaO CM velocities from reaction of Ba(¹P) (solid) and Ba(¹S) (dotted). Based on the increased BaO signal observed near the edges of the Newton sphere, we conclude that electronic excitation of the incident Ba leads to formation of BaO with slightly greater average translational energy than reaction of the ground state atom. Because the signal increased the same amount in the forward and backward hemispheres, formation of ground state BaO from electronically excited Ba primarily results from decay of long-lived collision complexes, as in the ground state reaction. Excitation of the barium atoms upstream of the collision zone results in 60% Ba (¹.³D), and 40% ¹S. The BaO angular distribution from Ba (¹.³D) was very similar; however, there was a smaller increase at wide lab angles since less energy was available from reaction of Ba(¹.³D) than Ba(¹P).

C. Ba + $NO_2 \rightarrow BaNO + O$

The occurrence of a chemically distinct second channel, BaNO + O was observed from Ba(¹S) + NO₂ at collision energies above 40 kcal/mole. The observed threshold collision energy from reaction of electronically excited Ba (^{1,3}D) was below 25 kcal/mole. Unlike BaO, the energy dependence of the BaNO fragmentation pattern cannot be easily obtained. Although this species is considerably more weakly bound than BaO, ionization of this radical likely involves removal of a nonbonding electron. This suggests that the internal energy dependence of the BaNO fragmentation pattern is probably not a major

concern. Indeed, the product time-of-flight for the NO⁺ daughter ion from BaNO could be observed and appeared to be the same as BaNO⁺. From the observation of the NO⁺ daughter ion, the location of the mass peak, and observation that the signal disappeared upon turning off the ionizer, we were able to rule out the possibilities that the signal resulted from BaO₂^{33a,b} leakage through the mass spectrometer, electronically excited or Rydberg Ba atoms⁴⁶, or nascent ions.³⁵

In the Ba(1 S) + NO₂ reaction, the BaNO product intensity was found to be rather weak, increasing with increasing collision energy. Unlike the dominant BaO + NO channel, BaNO from reaction of ground state Ba was only observed over a narrow range of laboratory angles, peaking away from the relative velocity vector. (Fig. 11, filled circles). The BaNO + O channel from Ba(1 S) + NO₂ appears to be completely independent of the BaO + NO channel. The BaNO time-of-flight spectra for reaction of Ba(1 S) are shown in Fig. 12. The slower broad peak in the time of flight spectrum remained even when the electron bombardment ionizer was turned off. It results from surface ionization of barium atoms at the back wall of the detector. The product angular and time-of-flight distributions were fit using a single uncoupled T(θ) and P(E), (Fig. 14). The BaNO product flux contour map from reaction of ground state Ba is shown in Fig. 15.

At a collision energy of 59 kcal/mole, electronic excitation of the Ba atom to the ^{1,3}D states resulted in a twofold increase in BaNO signal, and a larger increase was seen from the ¹P state. Unlike the ground state reaction, the best fit of the BaNO angular and velocity distribution-from-reaction of electronically excited Ba(¹P, ^{1,3}D) (Fig. 11- hollow

squares, Fig. 13) resulted in a forward-backward symmetric center-of-mass angular distribution (Fig. 14). The angular distribution obtained by pumping the Ba atoms upstream of the collision zone to the metastable ^{1,3}D states was weaker, but qualitatively similar. The forward-backward symmetric angular distributions indicate that unlike the BaNO resulting from reaction of ground state Ba(¹S), formation of BaNO from excited state Ba primarily results from decay of long-lived collision complexes.

IV. DISCUSSION

A. Ba +
$$NO_2 \rightarrow BaO + NO$$

i. Comparison to Past Work

Perhaps the most surprising result of our study is that the reaction Ba + NO₂ → BaO + NO primarily involves long-lived collision complexes with large releases of translational energy. Since both of these primary conclusions disagree with those from the past work of others, ^{18,19} we first address this discrepancy. The most obvious difference between this experiment and previous work is the higher collision energy in the present work. The past experiments were conducted using a single mean collision energy of ~2-3 kcal/mole whereas we studied the reaction at four energies with the lowest at 5 kcal/mole. An explanation for the discrepancy could be that a direct reaction mechanism with forward scattering occurs at the low collision energy of the previous work with the mechanism evolving into one with collision complexes at the higher energies employed here. This appears to be unlikely since the best fit to our data obtained at the lowest collision energy showed no enhancement of the forward scattering

component. Moreover, such behavior would be exactly opposite of what is usually observed:⁴¹⁻⁴⁴ a decreased statistical lifetime of the complex is normally expected with increasing energy, leading to direct reaction at higher energy.

Another possible explanation for our result is that at higher collision energies, the divalent barium atom preferentially inserts into a covalent bond with formation of O Ba²⁺NO. This would be somewhat analogous to the behavior we have recently observed in the reaction $Ba(^{1}D) + H_{2}O \rightarrow HBaOH^{\bullet} \rightarrow BaOH + H^{47}$, (Chapter 2). In such systems, a forward-backward angular distribution does not necessarily imply that the intermediates survive longer than one rotational period. Since insertion (or more likely, H-atom migration) could involve either of two identical O-H bonds, the symmetric BaOH angular distribution might simply reflect the C₂ symmetry of H₂O. Although it might seem that an analogous situation could exist here, we believe that such a situation is unlikely for two reasons. First, symmetry arguments indicate that insertion will be very unfavorable for Ba(1S) + NO₂. More importantly, insertion of ground state Ba(1S) into NO₂ is likely to involve a large potential energy barrier since the process involves a nearly simultaneous transition from a covalent to a doubly ionic surface. This suggests that a potential energy barrier will exist in the entrance channel. There should be a negligible barrier in the exit channel, however, since decay of the vibrationally activated insertion intermediate to BaO + NO should only involve simple bond rupture. Thus, we expect that the products from an insertion mechanism should be vibrationally excited with small translational energy. This is not consistent with the low vibrational excitation measured in the LIF experiments¹⁷ and large translational energy release that we observe. It

appears that an OBaNO insertion intermediate does not play an important role in the reaction.

We believe that the conclusions based on the previous crossed beams experiments were incorrect. The danger of making conclusions based on angular distributions alone has been addressed previously. Without velocity analysis, the transformation from the laboratory to center-of-mass angular distribution is not unique, especially when effusive beams are employed having broad velocity distributions. In particular, for a reaction of fast Ba atoms + slow NO₂ the transformation Jacobian will strongly enhance the forward peak, and the angular distribution alone will be rather insensitive to the actual $T(\theta)$ P(E) combination. Since the metal-oxide bond energies decrease for the lighter alkaline earth atoms, ²⁹ reactions of those species with NO₂ will be less exoergic. It is very likely that reaction of the other alkaline earths (Be is a possible exception) with NO₂ will also involve complex intermediates. ⁴⁹

ii. Mechanism for Ba + NO₂ → BaO + NO

There is no question that this reaction is initiated by the well known "harpoon mechanism" 2,3 used to describe reactions such as $K + Br_2 \rightarrow KBr + Br$. The energies of the lowest electronic states of NO_2^{-50} are summarized in Table III together with the calculated ionic-covalent curve crossing radii and estimated reaction cross sections. The very large experimental reaction cross section can only be consistent with long range transfer into the half filled $6a_1$ orbital (Fig. 16), corresponding to formation of ground state $NO_2^{-(1}A_1)^{50}$. The analogy between the present reaction and the alkali-halogen

reaction extends no further, however. In the case of bromine, not only is the process $Br_2 + e \rightarrow Br_2 \rightarrow Br + Br$ exoergic, ²⁹ but vertical electron attachment to Br_2 results in formation of Br_2 on the inner repulsive wall of the Br-Br interaction potential at an energy near the threshold for dissociation. ⁶ Electron attachment to NO_2 , however, results in formation of a strongly bound anion; in fact $D_o(O^-NO)$ actually exceeds $D_o(O^-NO)$. In low energy electron scattering, it is not surprising that the process $Br_2 + e^- \rightarrow Br^- + Br$ is observed at electron energies down to $O(V^{-1})$. The energy threshold for $NO_2 + e^- \rightarrow O^- + NO$, however, is equal to its endoergicity, 1.61eV. ⁵²

Although there is no information available as to the structure of gaseous BaNO₂, it was noted previously that considerable similarity is expected to the analogous alkali nitrites which are known to be stable molecules, even in the gas phase.⁵³ We have also noted that an OBaNO structure is unlikely. There appears to be no question that the alkali nitrites are ring shaped C_{2v} molecules with the metal cation symmetrically located in the plane of NO₂. This structure is supported by spectroscopic measurements both in matrix isolation experiments⁵⁴ and in the gas phase.⁵⁵ Ab initio calculations also indicate that a ring shaped structure is of lowest energy.⁵⁶ Experimental results on the alkaline earth nitrites⁵⁷ in rare gas matrices suggest the same structure or possibly a nonplanar structure with a poorly defined position of the M⁺ above the plane of the anion.

Orbital symmetry considerations (Fig. 16) indicate that interaction of Ba⁺ with $NO_2^- 6a_1^2$ (1A_1) should be particularly favorable in the planar configuration in which the Ba⁺ lies on the C_{2v} axis between the two oxygen atoms. This will maximize the ionic attraction since the electron density on NO_2^- can be delocalized over the oxygen atoms

and at the same time interact strongly with Ba^+ . In addition, a net covalent interaction can be achieved by favorable overlap of the $Ba^+(6s)$ orbital and the $NO_2^ 6a_1$ orbital. Other planar geometries (such as C_a) should be less stable due to poor orbital overlap. The two higher lying excited states of NO_2^- result from transfer of an electron into the unfilled $2b_1$ orbital leading to the 1B_1 and 3B_1 states. Not only does this orbital lie \sim 3eV higher than the $6a_1$ (Table III), but based on orbital symmetry it appears that interaction of the excited states of NO_2^- with Ba^+ will be weaker than for the ground state since overlap of the orbitals lead to no net covalent attraction.

For nearly all trajectories, long range electron transfer into the $6a_1$ orbital should be very favorable at $r \approx 4.9 \text{Å}$. Based on the geometries for NO_2 and NO_2 (Table IV), and the O-O bonding character of the $6a_1$ orbital, electron transfer will result in symmetric bending of NO_2 with only a small amount of NO vibrational excitation.²² We note that the asymmetric stretching mode (formally corresponding to the reaction coordinate) is not expected to be appreciably excited. Approach of Ba directly towards N along the C_{2v} axis is also expected to lead to efficient electron transfer but the resulting C_{2v} BaNO₂ molecule is probably less stable than the bidentate C_{2v} structure. However, such a species is still likely to survive for at least several vibrational periods since the departing O ion will be initially moving along one N-O bond axis. For a monodentate C_{2v} Ba⁺-NO₂ molecule, this motion will oppose the Ba⁺-O coulombic attraction, leading to a longer lifetime for decay of the ionic intermediate.

This reaction is very unusual since it is dominated by a mechanism involving collision complexes despite the fact that a very highly exoergic decay channel (BaO +

NO) is known to be accessed with a probability of nearly 100%. Since it is known that the slightly endoergic reaction Cs + NO₂ → CsO + NO proceeds readily at a collision energy of 3 kcal/mole, 25 the potential energy barrier for decay of Cs+NO₂ to products must be nearly zero. If the barrier for decay of Ba+NO₂ to Ba²⁺O²⁻ + NO was also negligible, survival of long-lived complexes for $\tau \ge 1$ psec would require a Ba-NO₂ binding energy of >250 kcal/mole.⁵⁸ This is far larger than is reasonable. We attribute the existence of complexes and the large amount of product translational energy from their decay to a large exit potential energy barrier for formation of products. This barrier in the exit channel appears to be a consequence of the divalent character of Ba. Decay of the complex to ground state molecules requires a substantial rearrangement of electronic structure, corresponding to transfer of the second valence electron necessary for formation of Ba²⁺O². The large exit barrier for decay of the complex apparently results from several factors. One factor is related to the most stable geometry for Ba⁺NO₂ likely being planar. Although the NO₂ energy levels will be shifted significantly by the presence of Ba⁺, it appears that the electronic rearrangement necessary for decay to products requires transfer of electron density into the 2b, lowest unoccupied molecular orbital of NO₂. As can be seen from Fig. 14, this is forbidden by orbital symmetry constraints for planar Ba⁺NO₂ complexes.

The height of the exit barrier for electronic rearrangement can be estimated from this work. It must lie below the energy of the reactants since we know that most complexes decay to products rather than back to reactants. Simple RRKM calculations⁵⁸ using $D_o(Ba-NO_2)=65$ kcal/mole and reasonable estimates of BaNO₂ vibrational

frequencies suggest that at collision energies above 30 kcal/mole, complex lifetimes of ≥ 1psec will result only if the barrier for decay of BaNO₂ is greater than ~45 kcal/mole. Using a 65 kcal/mole well depth, this implies that the barrier is not lower than 20 kcal/mole below the energy of separated Ba + NO₂. Based on our observation that the observed Ba⁺ signal primarily results from fragmentation of BaO in the ionizer, (rather than from decay of BaNO₂ back to Ba + NO₂), we believe that our lower limit of 45 kcal/mole is likely to be close to the true value. If the barrier were much nearer to the energy of separated Ba+ NO₂, nonreactive scattering from decay of BaNO₂ back to reactants should lead to an additional component in the Ba time-of-flight spectrum with relatively low translational energy release. Figure 17 shows the energy levels of all relevant chemical species involving ground state Ba-- solid lines denote reaction pathways.

B. Source of the Forward Scattered BaO Channel

Our experimental data indicates the existence of a second reaction channel leading to forward scattered BaO with smaller translational energy release than from the dominant channel. This channel results from a direct reaction mechanism; based on the observed product translational energy distribution, by conservation of energy the reaction exoergicity must be primarily channeled into product internal excitation. It might seem plausible that this direct reaction channel corresponds to formation of electronically excited BaO, best described as Ba+O·.30,59 This explanation would appear to be particularly appealing since second electron transfer is not required for this product.

However, our measurements indicate that the forward scattered channel represents $20\pm5\%$ of the total reaction cross section whereas the single collision chemiluminescence yield is known to be less than $0.2\%.^{16,17}$ Since there is no reason to believe that electronically excited products should be preferentially formed in dark states, assuming that the chemiluminescence yield is correct, a large fraction of the forward scattered BaO products must be vibrationally excited ground state molecules.

Since a direct reaction mechanism implies a short lifetime for the ionic intermediate, it might seem at first surprising that the direct reaction mechanism is relatively minor. This indicates that the forward scattered BaO does not result from kinetic competition with the dominant channel involving collision complexes. Instead, the relatively small yield of direct, forward scattered products must reflect a small probability for avoiding the deep Ba⁺NO₂ well. There are two possible mechanisms whereby this may be achieved. Some trajectories can remain on the covalent surface by avoiding electron transfer at the first crossing. In this case, electron transfer at the second crossing is nearly certain and the reaction intermediate can be considered to be an excited state of Ba⁺NO₂ which will have a shorter lifetime due to its weaker bond. This mechanism appears to be dominant in the reaction Li + NO₂ \rightarrow LiO + NO, which was found to be direct without the involvement of long-lived collision complexes.²⁷ This model appears unlikely in the present case, however, since the relative yield of forward scattered products is not very sensitive to collision energy. Since adiabatic behavior at the outer crossing is expected to decrease with increasing collision energies, 60 if this explanation were correct we should have observed a substantial increase of forward

scattering with increasing collision energy. Moreover, correlations⁶¹ between curve crossing distance and electron transfer probability in atom-atom and atom-diatom systems indicate that at 4.9Å, the electron transfer probability will be nearly 100%. The alternative explanation is that the direct reaction results from electron transfer into the 6a₁ orbital from Ba-NO₂ geometries which are unfavorable for formation of long-lived complexes. This situation would be expected to occur in those collisions where the Ba⁺ can approach very close to one oxygen atom in NO₂. In this case, a very close Ba⁺-O interaction would be favorable for second electron transfer and formation of ground state products. Good coupling between the incident kinetic energy and the reaction coordinate facilitates second electron transfer and formation of vibrationally hot, forward scattered BaO.

C. Ba +
$$NO_2 \rightarrow BaNO + O$$

i) The Ba(1S) Reaction Mechanism

A number of observations allow us to gain considerable insight into the dynamics of this channel. The angular distribution from reaction of ground state Ba(1S) was found to peak over a very narrow range, well away from the relative velocity vector. Formation of BaNO results from a direct reaction, unlike the dominant ground state BaO channel. The small cross section and a narrow angular distribution indicates that successful reaction results from a very restricted range of collision geometries.

In the next section, we will argue that BaNO is best represented as Ba+NO. An

obvious question is whether long range electron transfer into the 6a₁ orbital of NO₂ precedes formation of BaNO + O. We believe that it does not; rather, BaNO likely results from trajectories which are able to avoid harpooning at the outer crossing point. The probability density maps for electrons in the 6a₁ and 2b₁ orbitals (Fig. 16) are useful in determining favorable and unfavorable molecular orientations for electron transfer. Although the shapes of the orbitals will become distorted upon close approach of Ba, they should be reasonable approximations outside the first ionic-covalent curve crossing radius (4.9Å). Transfer of an electron from the spherically symmetric Ba 6s orbital into the NO₂ 6a₁ orbital can be best avoided by approach of Ba towards N just off the C_{2v} axis. This approach geometry will also lead to asymmetric stretching of NO₂, an important requisite for reaction. The substantial threshold and narrow product angular spread is consistent with a narrow approach geometry and a need to couple translational energy into the asymmetric stretching of NO₂. However, maintaining proper collision geometry alone is not likely to be sufficient to promote this reaction. Since electron transfer into the 6a₁ orbital leads to symmetric bending or symmetric stretching, only by remaining on the covalent surface at the outer crossing is it possible to excite the asymmetric NO₂ motion. The small product yield (<2% that of BaO) is consistent with a very small probability for avoiding electron transfer at the first ionic-covalent crossing.

Based on the translational energy measurements, we calculate $D_o(Ba-NO) \ge 65$ kcal/mole (Section iii). Since $D_o(O-NO)=72$ kcal/mole²⁹, the reaction is only slightly endoergic. However, the energy threshold for this reaction from Ba(1S) appears to be

much greater than the endoergicity. It appears that the threshold is primarily due to a dynamic rather than thermodynamic requirement. The dynamic requirement is probably the need for nonadiabatic behavior at the outer crossing point and good coupling of the incident kinetic energy into the reaction coordinate, i.e. asymmetric N-O stretching. This is supported by our observation that the reaction is strongly promoted by very high collision energy. The probability of long range electron transfer depends on several factors. 60,61 These include the difference in the gradients for each potential energy surface in the vicinity of the crossing region, impact parameter, orientation of the NO, molecule, radial velocity, and coupling matrix element. For the reaction of ground state Ba(1S), formation of BaNO requires small impact parameter collisions. For small impact parameters, the relative velocity is almost entirely radial and an increase in collision energy can lead to substantially decreased adiabatic behavior and asymmetric motion of the NO₂ molecule. This should lead to enhancement of BaNO by decreasing the probability for outer harpooning and permitting good coupling of the incident kinetic energy into the reaction coordinate.

ii) The Effect of Electronic Excitation

Electronic excitation to the ¹P state increases the first ionic-covalent curve crossing radius to ~15 Å, leading to a substantially decreased probability for electron transfer into the 6a₁ orbital.⁶¹ The next crossings correspond to transfer of the Ba 6s electron into the 6a₁ orbital (retaining the Ba 6p core) or transfer of the 6p electron into

the 2b₁ orbital. Both processes correspond to formation of excited state Ba⁺NO₂. Based on the BaO angular distributions, a large fraction of these excited complexes still survive for longer than one rotational period. At least part of this increased energy appears in relative translational motion of the recoiling BaO/NO products.

With electronic excitation to the ^{1,3}D states, we observed an approximately twofold enhancement of formation of BaNO + O. The yield was larger from the (¹P) state with an enhancement by a factor of 3-4. However, the products appear to result from decay of long-lived complexes, whereas the reaction is direct from the ground state. This unusual behavior likely results from a substantial increase in the energy of the Ba⁺NO₂ complex due to electronic excitation of the incident Ba atom. In the case of ground state Ba(¹S), only a very small fraction of collisions could avoid the first electron transfer. With proper collision geometry and good coupling of incident collision energy into the reaction coordinate, BaNO + O resulted from a direct reaction mechanism. In the reaction of electronically excited Ba, decay of internally excited Ba⁺NO₂ complexes leads to the formation of BaNO + O products in addition to the dominant BaO + NO.

iii) The Dissociation Energy of BaNO

The Ba-NO binding energy may be derived from the translational energy distribution for Ba(1 S) + NO₂ \rightarrow BaNO + O. The following expression relating the energy of the reactants to that of the products is based on conservation of energy:

$$E_{coll} + E_{int,NO2} - D_o(O-NO_2) = E_{int,BaNO} + E_{int,O} + E_{trans,BaNO/O} - D_o(Ba-NO).$$
 (3)

By assuming efficient cooling of rotational but not vibrational energy of NO_2 in the supersonic expansion, $E_{int, NO2} \approx 2$ kcal/mole. The following equation, derived from (2) is actually an equality if the fastest BaNO products correspond to formation of BaNO in its ground rovibrational states and the oxygen atom is $O(^3P)$:

$$D_o(Ba-NO) \ge E_{trans, BaNO/O} + D_o(O-NO_2) - E_{coll} - E_{int, NO}$$
 (4)

Using $E_{coll} = 60$ kcal/mole and $D_o(O-NO_2) = 72$ kcal/mole, we calculate $D_o(Ba-NO) \ge 65$ kcal/mole, with the uncertainty not greater than 20 kcal/mole. Uncertainties arise in fitting the translational energy distribution and from the spread of the velocity distribution of the Ba beam. Since it is likely that at least some ground state products are formed in the reaction, we conclude that $D_o(Ba-NO) = 65 \pm 20$ kcal/mole.

There has been no previous direct observation of BaNO in the gas phase. In fact, as far as we know, this is the first direct observation of an isolated gaseous metal mononitroxide. However, infrared spectra of alkali⁶² and alkaline earth⁵⁷ nitroxides have been recorded under matrix isolation conditions. Based on that work, it was concluded that the molecules are ionic: isotope shifts in the IR spectra suggested a bent or linear M⁺NO molecule (rather than M⁺ON) in all cases except Li⁺ON. Although no binding energies are available for other nitroxides, it is interesting to compare BaNO to known

metal-dioxygen compounds. As perhaps one might expect, $D_o(Ba-NO)$ is comparable to singly ionic metal superoxides rather than the alkaline earth-dioxygen molecule which appears to have substantial doubly ionic character: $D_o(Na-O_2) = 58.1 \pm 2 \text{ kcal/mole}^{63}$, $D_o(Ba-O_2) = 120 \pm 20 \text{ kcal/mole}^{33a,b}$.

D. The Absence of Ba + $NO_2 \rightarrow BaO_2 + N$

Considerable effort was devoted to observing this reaction since there was good reason to believe that it might occur. Electron transfer into the $6a_1$ orbital strongly enhances bonding between the oxygen atoms with the O-N-O bond angle decreasing from 134.1° to 115.4° (Table IV). The reaction $B + NO_2 \rightarrow BO_2 + N$ has been observed by Green and $Gole^{64}$ and was attributed to rapid scissoring of the NO_2^- molecule after electron transfer. Other triatomic oxidants such as O_3 and ClO_2 did not lead to formation of BO_2 since the bond angles in those molecules are not appreciably different from their anions. Dissociative electron attachment to NO_2 is known to lead to formation of O_7 , NO^- and O_2^- at 1.61 eV, 3.11 eV, and 4.03 eV, respectively. Since these energies are the thermodynamic thresholds for the processes, $O_2^{0.52}$ we postulated that with electronic excitation or at high collision energy, nascent O_2^- might be formed from reaction of O_2^- at O_2^- might be formed from reaction of O_2^- at O_2^- might be formed from reaction of O_2^- at O_2^- might be formed from reaction of O_2^- at O_2^- might be formed from reaction of O_2^- might be formed from reaction of O_2^- might O_2^- might be formed from reaction of O_2^- might O_2^- might be formed from reaction of O_2^- might O_2^- might be formed from reaction of O_2^- might O_2^- migh

The BaO_2 + N channel either does not occur at all, or is below the sensitivity of the experiment. There was no evidence that electronic excitation promotes this channel. The cross section must be below $1\dot{A}^2$ for formation of BaO_2 + N. It appears that the

contrasting behavior between B and Ba arises primarily due to the ionization potential of B $(8.3\text{eV})^{21}$ being considerably higher than that of Ba $(5.2\text{eV})^{21}$. In the case of Ba, electron transfer occurs at long range ($\sim 4.9\text{Å}$) whereas much closer distance is required for B ($\sim 2.4\text{Å}$). This difference appears to be critically important to the outcome of the chemical reaction. In the case of B, electron transfer and symmetric bending of NO₂ will only occur when the B atom is essentially within the attractive well of the very strongly bound B-O₂ interaction potential. In the case of Ba, however, the ionic complex will execute many vibrations before the Ba⁺ can approach within the close range required for formation of BaO₂. Intramolecular energy redistribution within the complex leads to randomization of the initial symmetric bending motion. This results in long-lived complexes which decay to BaO + NO-- apparently the potential energy barrier for formation of BaO₂ + N is large. In those collisions where electron transfer into the 6a₁ orbital can be avoided, the dominant chemical product is BaNO + O rather than BaO₂ + N.

V. CONCLUSIONS

We have found the chemistry of the Ba + NO₂ reaction to be substantially more complex than previously believed. Our results on Ba + NO₂ disagree with the earlier conclusions, ¹⁸ which were based solely on product angular distributions. We conclude that long-lived Ba⁺NO₂ intermediates are involved in the majority of all reactive trajectories despite reaction exoergicities that are comparable to complex well depths.

This rather unusual behavior appears to be a dynamic consequence of the divalent nature of barium: decay of the initially formed complex to ground state molecules involves a substantial potential energy barrier. This barrier is associated with the electronic rearrangement necessary for the initially formed singly ionic Ba⁺NO₂ complex to decay to ground state doubly ionic Ba²⁺O²⁻+ NO. A very large release of translational energy, the magnitude of which is perhaps unprecedented from decay of collision complexes, results from a barrier to second electron transfer followed by strong repulsion of the stable BaO + NO products.

Two additional direct reaction channels were observed. A minor BaO + NO channel led to forward scattering of the metal oxide with a relatively small release of translational energy. Our observation that the yield of forward scattered products is larger than the known chemiluminescent yield indicates that this channel corresponds primarily to formation of vibrationally hot ground state BaO from direct reaction involving small impact parameter collisions. A previously unknown molecule, BaNO, was observed as a nascent minor product from Ba + NO₂. Unlike the dominant BaO channels that involve outer harpooning, formation of BaNO from Ba(¹S) likely results from the small fraction of collision geometries that avoid long range electron transfer.

Even with electronic excitation of the incident barium atom, reactions involving long-lived collision complexes continued to dominate the neutral product channels.³⁵ Electronic excitation results in increased internal excitation of the Ba⁺NO₂, leading to a larger average translational energy release in the BaO + NO products. Whereas

complexes resulting from collisions of ground state Ba only decayed to BaO + NO, a small fraction of those from electronically excited Ba also decayed to BaNO + O.

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TABLE I. Thermodynamics of Possible Ba(1S) + NO₂ Reactions

Reaction	ΔH (kcal/mole)	
$BaO(X^1\Sigma^+) + NO$	-61ª	
$BaO(A^1\Sigma^+) + NO$	-11*	
$BaO(A'^{1}\Pi,a^{3}\Pi) + NO$	-11*	
BaNO + O	+7 ^b	•
$BaO_2 + N$	-12°	

Reference 29. BReference 33a References 33a and 33b

TABLE II. Measured Atomic and Molecular Beam Velocities and Speed Ratios

Beam	Seed Gas (Pa)	Beam Velocity (m/s)	Speed Ratiob
Ba	Kr (300)	860	12
Ва	Ar (400)	1200	14
Ba	He:Ne ^c (550)	1960	14
Ba	He (700)	3560	16
$NO_2 \sim 10\%$	He (250)	1520	14
NO ₂ ~ 20%	He (250)	1210	12
NO ₂ (100%)	None	720	6

^aPressure in Torr ^bv/Δv FWHM. ^c1:1 Mixture

TABLE III. States of NO₂

State		E(eV)ª	$R_x(A)^b$	$\sigma_{\rm calc}({\rm \AA}^2)^{\rm c}$
¹ A ₁	6a ₁ ²	-2.27 ^d	4.9	75
3B_1	$6a_1^{1},2b_1^{1}$	+0.57°	2.5	20
¹ B ₁	$6a_1^1,2b_1^1$	+1.2°	2.2	15

- a. Energy relative to $NO_2 + e^-$.
- b. Ionic covalent curve crossing radius calculated using $R_x(A) = 14.4/(I.P._{Ba}-E)(eV)$.
- c. Estimated harpooning cross section based on $\sigma_{calc} = \pi R_x^2$
- d. Electron affinity of NO₂ from Ref 22.
- e. Energies relative to $NO_2 + e^-$ based on NO_2 electron affinity and excited state energies tabulated in Ref 50.

TABLE IV: Molecular Parameters

Species	R _{N-O} (Å)	α (degrees)	
$NO_2(X^2A_1)$	1.19389(4) ^a	133.857(3)ª	
$NO_2^-(X^1A_1)$	1.25(2) ^a	$117.5 \pm 2.^{a}$	
NO(X ² II)	1.1508 ^b	-	

a. Ref 22, and references therein.

b. Ref 65

VIII. FIGURE CAPTIONS

- 1. Top view of the crossed molecular beams apparatus.
- 2. BaO laboratory angular distribution for Ba + NO₂ → BaO + NO at collision energy of 12.5 kcal/mole. o Experimental points. Total fit to data. …… Contribution from channel 1. Contribution from channel 2. The nominal Newton diagram is shown with a limiting circle indicating the maximum center-of-mass velocity for BaO based on the known reaction exoergicity for formation of ground state products.
- BaO product time-of-flight spectra at indicated laboratory angles for Ba + NO₂
 → BaO + NO at collision energy of 12.5 kcal/mole. o Experimental data.
 Total fit to data.
 - Contribution from channel 1. ----- Contribution from channel 2.
- 4. Best fit translational energy P(E) and center-of-mass angular distribution T(θ) for the BaO + NO products. Collision energy was 12.5 kcal/mole. Channel 1 represents the channel involving long-lived BaNO₂ collision complexes and channel 2 is the direct forward scattered channel.
- 5. BaO product angular distributions obtained at mean collision energies of 31 and 59 kcal/mole.
- 6. Same as Fig. 3 but $E_{coll} = 59$ kcal/mole.
- 7. Same as Fig. 4 but $E_{coll} = 59$ kcal/mole.

- 8. BaO product flux contour map at $E_{coll} = 12.5$ kcal/mole. A- Newton diagram B-Contribution from channel 1. C- Contribution from channel 2. C- Total BaO product flux (Channel 1 + Channel 2).
- 9. BaO product flux contour map for channel 1 + channel 2 superimposed on the nominal Newton diagram. $E_{coll} = 12.5 \text{ kcal/mole}$.
- 10. BaO product angular distribution at collision energy of 12.5 kcal/mole showing effect of electronic excitation of the incident Ba atoms. Laser excitation was at the collision zone. Beam consists of approx. 30% ^{1,3}D, 35% ¹P, 35% ¹S. Nominal Newton diagram showing circles representing maximum BaO velocities from reaction of ground state Ba(¹S)(dotted line) and electronically excited Ba(¹P) (solid line).
- 11. BaNO product angular distribution at collision energy of 59 kcal/mole. Laser off:

 O-Experimental points, --- Fit to data. Laser focussed at interaction region: □
 Experimental points, --- Fit to data. Newton diagram is also shown.
- 12. BaNO product time-of-flight data from reaction of Ba(1 S) at indicated laboratory angles for Ba + NO₂ \rightarrow BaNO + O. Collision energy was 59 kcal/mole. o-Experimental points. Fit to data.
- 13. Same as Fig. 12 but with laser excitation at interaction region.
- Best fit translational energy P(E) and center-of-mass angular distribution T(θ) for the BaNO + O products. Collision energy was 59 kcal/mole. --- Laser off.
 Laser excitation at interaction region.

- 15. Center-of-mass product flux contour map in velocity space for the BaNO product from the Ba + NO_2 reaction. Laser is off. $E_{coll} = 59$ kcal/mole.
- 16. Electron probability density maps for the half occupied $6a_1$ and unoccupied $2b_1$ orbitals of NO_2 .
- 17. Energy level diagram for reaction of Ba(1S) + NO₂. Shaded area indicates uncertainties in thermodynamic quantities.

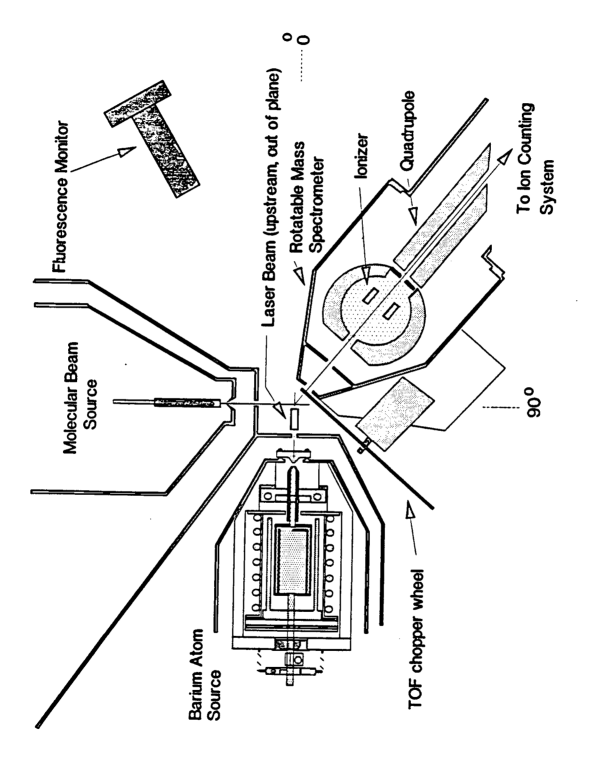


Fig. 1

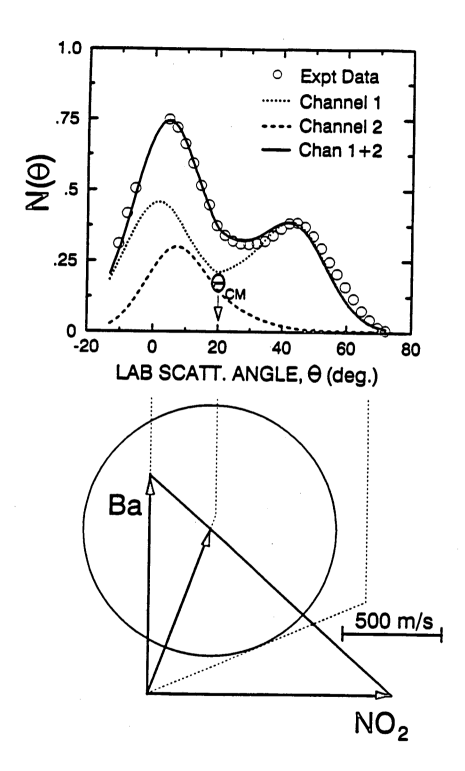


Fig. 2

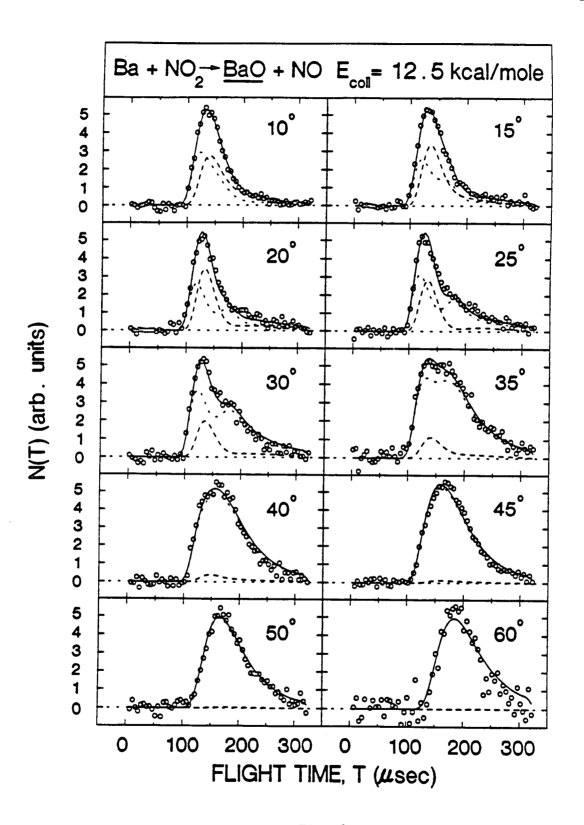


Fig. 3

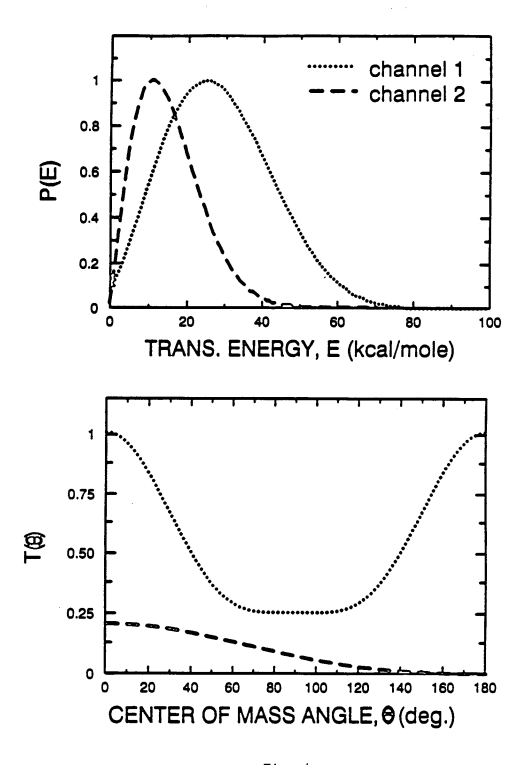


Fig. 4

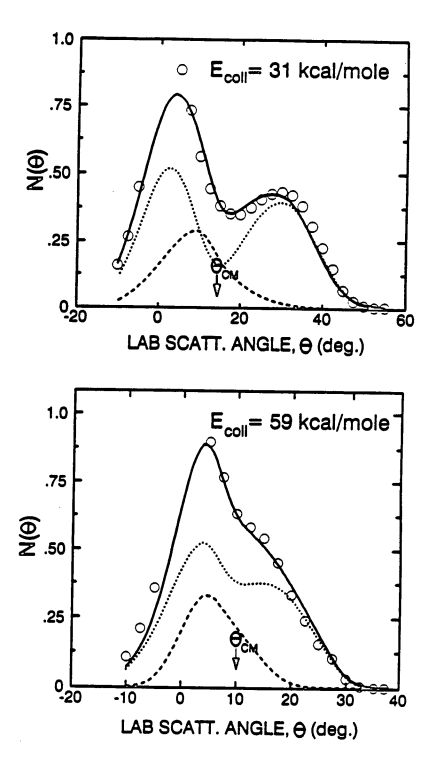
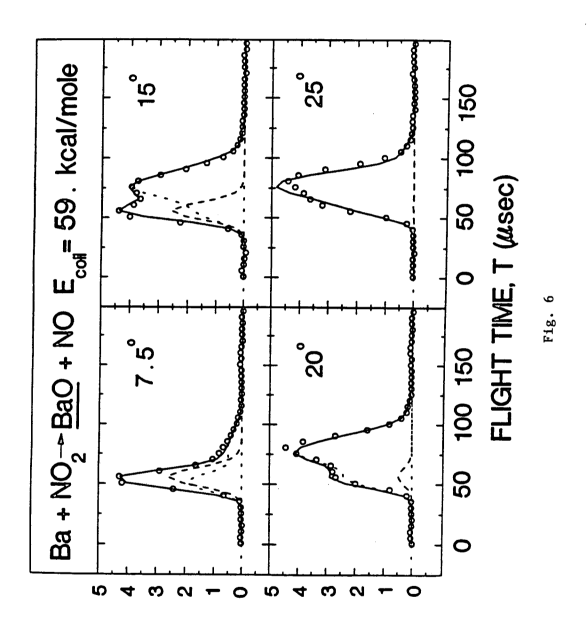


Fig. 5



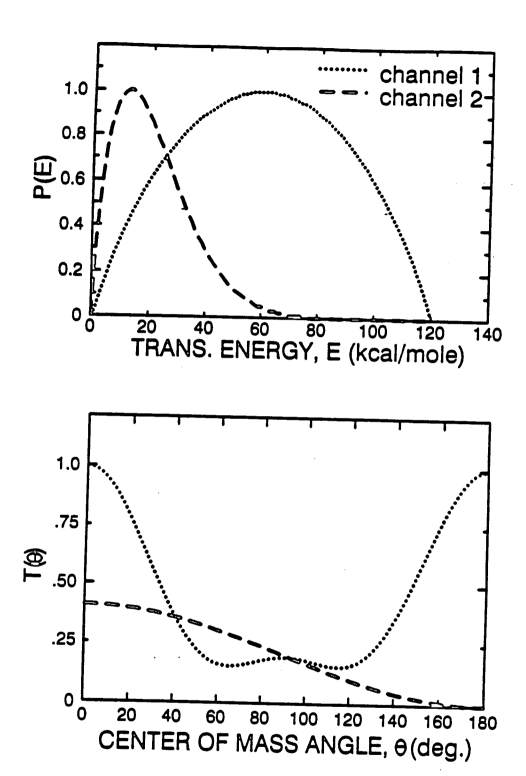


Fig. 7

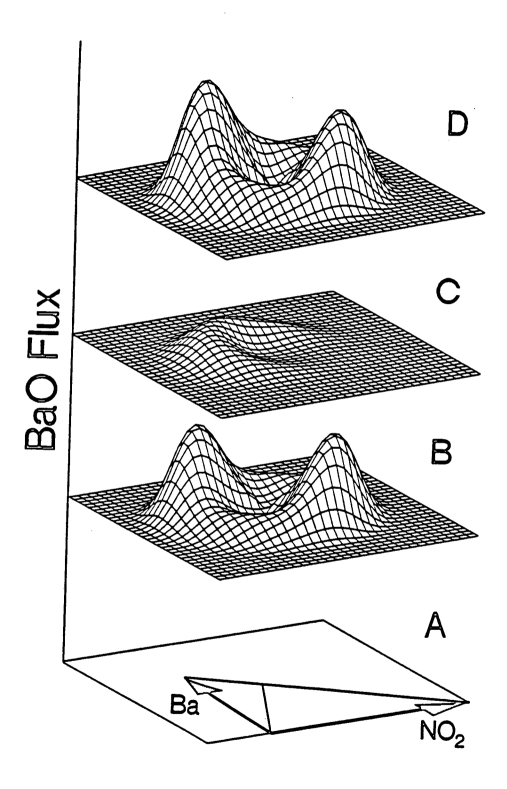
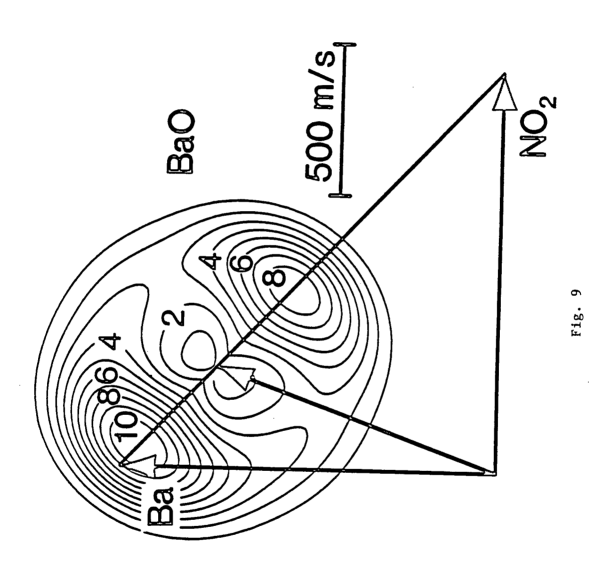


Fig. 8



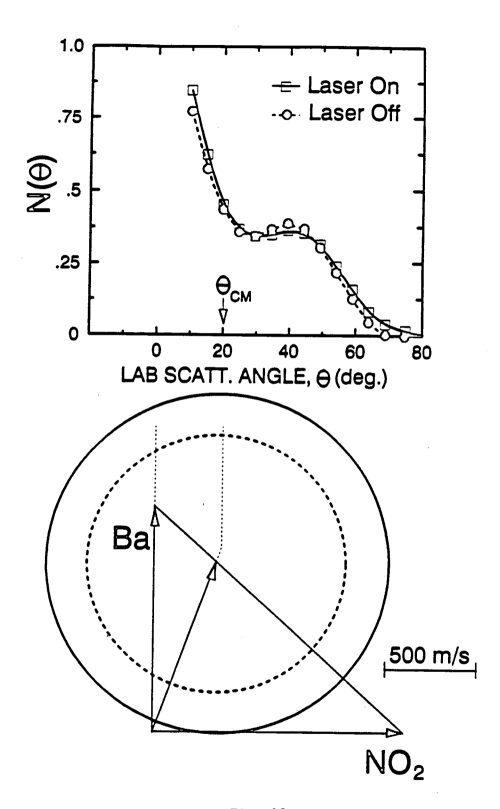


Fig. 10

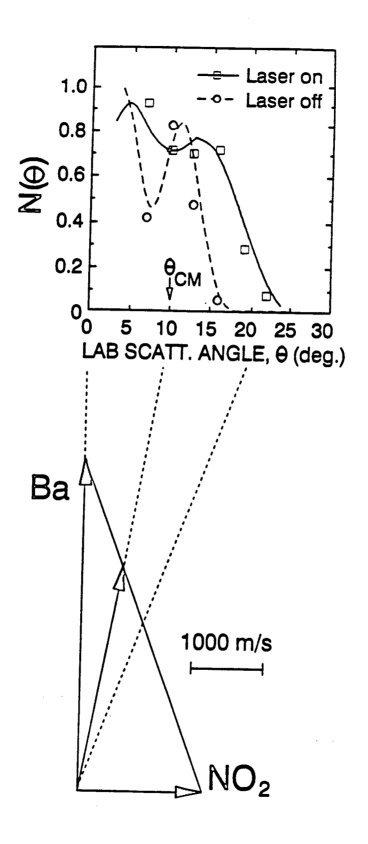


Fig. 11

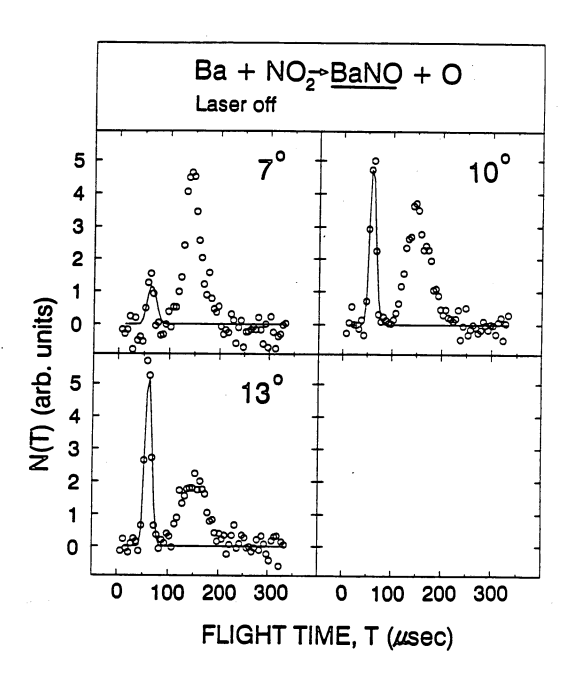


Fig. 12

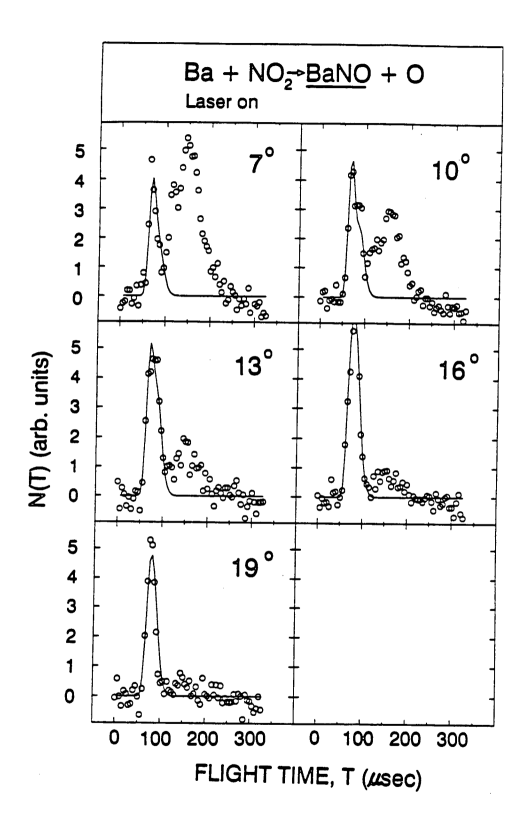
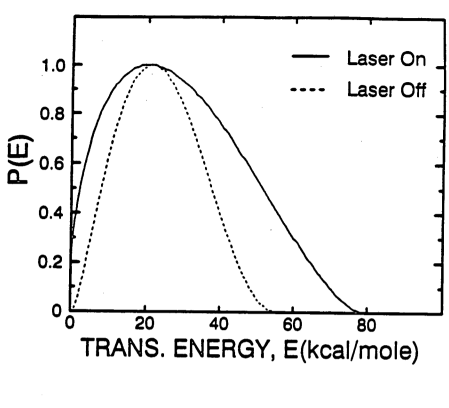


Fig. 13



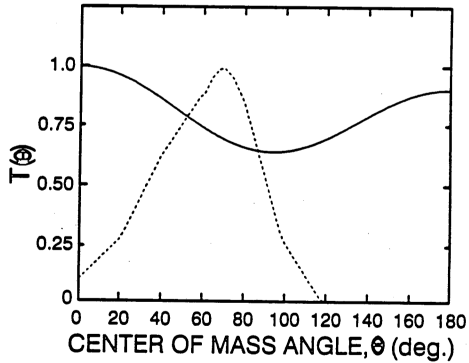


Fig. 14

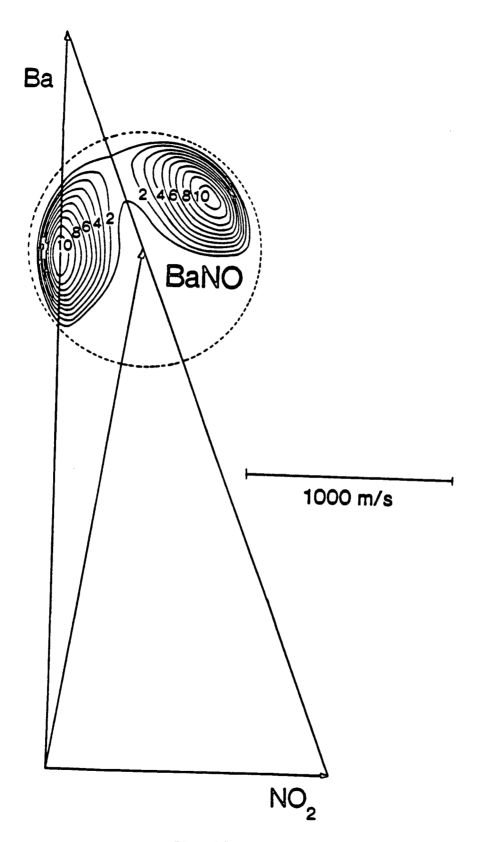


Fig. 15

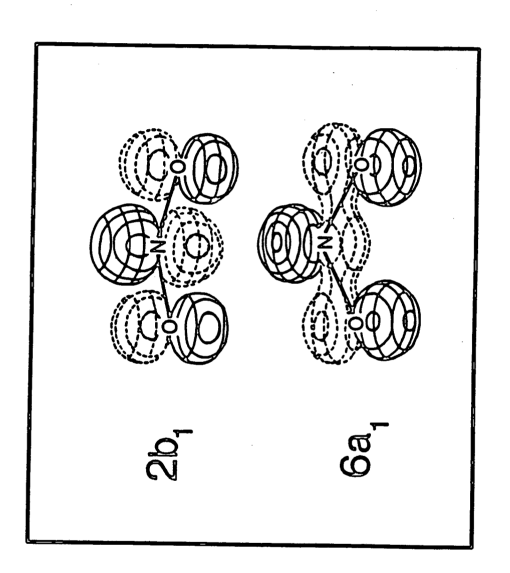
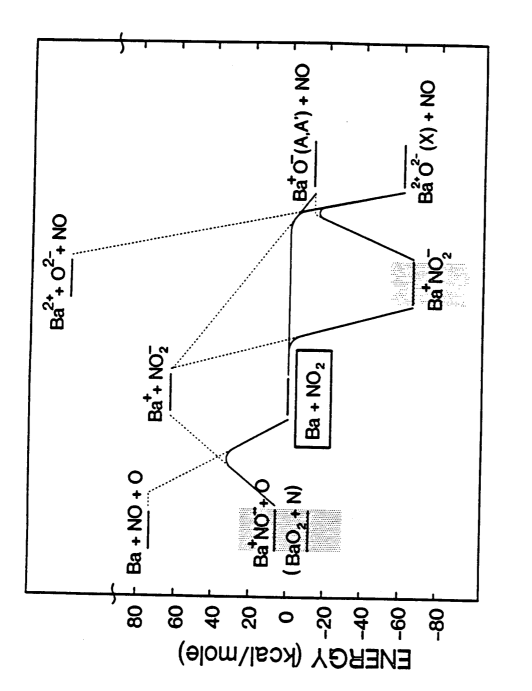


Fig. 16



1g. 1

CHAPTER 2

State Specific Reactions of Ba(¹S₀) and Ba(¹D₂)
With Water and its Alkyl Derivatives.

ABSTRACT

The reactions of Ba(6s²-1S₀) and Ba(6s5d-1D₂) with water and alcohols were studied under single collision conditions using crossed molecular beams. Reaction of ground state Ba(1 S) + H₂O led to dominant formation of BaO($X^{1}\Sigma^{+}$) + H₂ at all collision energies studied but the formation of BaOH + H could be observed weakly at collision energies above the endoergicity of the reaction (13 kcal/mole). Electronic excitation of the Ba atom reactant to the metastable ¹D state led to large enhancement in formation of BaOH + H but not BaO + H_2 . The dominance of BaO + H_2 from ground state Ba(1 S) even at collision energies nearly equal to the Ba(¹D) excitation energy (32kcal/mole) indicates that the observed reaction state specificity results from participation of different potential energy surfaces for reactions of ground and excited state atoms. Collisions of Ba(1S) and Ba(1D) with methanol led only to formation of BaOCH₃ + H, whereas reaction with allyl alcohol (H₂C=CH-CH₂OH) resulted in formation of both BaOCH₂CH=CH₂ + H and BaOH + CH₂=CH-CH₂. Large enhancements in reactivity with all alcohols were observed upon electronic excitation but no reaction was observed from Ba(1S) or Ba(1D) + dimethyl ether (CH3OCH3). This behavior strongly suggests that the reactions leading to radical formation are dominated by H-atom migration.

I. INTRODUCTION

By judicious choice of the electronic state of a reactant it is possible to affect or even control the chemical outcome of a bimolecular reaction. Recent work¹ using guided ion beams has shown that the chemistry of the various electronic states of a transition metal ion can change markedly due to the participation of different potential energy surfaces. In the case of neutral species, early studies showed that O(¹D) is able to insert into covalent bonds, whereas the ground state O(³P) atom is relatively inert under the same conditions.² More recently, the use of crossed supersonic atomic and molecular beams combined with laser excitation of the atomic reactant has made it possible to study the reaction dynamics of different electronic states of an atom as a function of collision energy under well defined conditions.³⁻⁵

Electronic excitation does not simply provide energy to drive endoergic reactions. Instead, the differing topologies of the potential energy surfaces will determine the reactivities of different states. This was demonstrated in the reaction $Na^* + O_2 \rightarrow NaO + O.^{3,4}$ When excited to the 4d state, sodium was found to react strongly under single collision conditions, whereas the 5s state (at nearly the same electronic energy) was unreactive.

There has been considerable interest in the interactions between water and isolated metal and main group atoms in the ground and electronically excited states. 6-15 In addition to being of importance in understanding the mechanisms of hydration, these interactions are intriguing because, unlike most small oxygen-containing molecules, water and its methyl derivatives posses only "Rydberg- like" unfilled molecular orbitals. 16,17

Consequently, reactions are not expected to be initiated by long range electron transfer, and the well known "harpoon mechanism" ^{5a,18-20} is not generally applicable. Instead, a close collision is necessary to initiate a chemical reaction between the atom and molecule. Because Ba is divalent, the chemistry of these systems is quite rich with several competing channels possible. In the case of water or methanol, insertion of the Ba atom into a covalent bond may occur, possibly forming HBaOH, HBaOCH₃ or HOBaCH₃^{6,7}. Decomposition of these intermediates would likely lead to formation of radical products BaOH or BaOCH₃. Alternatively, abstraction of the central oxygen atom resulting in formation of the closed shell molecules BaO + H₂ or CH₄ is actually the thermodynamically favored channel. ²¹(Table 1)

The reactions of alkaline earth and a variety of other atoms (M) with water have been studied extensively by Margrave and coworkers ^{7,8,14,15} using matrix isolation techniques and infrared spectroscopy. They found that codeposition of the reactants in an argon matrix led to formation of a M-OH₂ adduct involving a weak interaction (5-10 kcal/mole) between the metal atom and oxygen in water (Table 1). An increasing red shift in the water bending frequency with the heavier alkaline earth atoms has been taken as evidence that the interaction is strongest for barium^{7,9}. For alkaline earth and some transition metal atoms, irradiation of the matrix at wavelengths near the metal atomic resonance line led to formation of HMOH and MOH. However, in the case of some atoms, the oxygen bonded adduct rearranged spontaneously in the dark to form HMOH. Recent ab initio calculations^{9,10} for the lighter alkaline earth atoms + water were consistent with these observations. The oxygen bonded adduct was found to represent

a small but definite well on the potential energy surface; a substantial barrier for insertion explained the need for irradiation to promote the reaction. Based on ab-initio calculations⁹ and empirical bond energy estimates⁷, HMOH is expected to be a stable molecule, lying about 30 kcal/mole lower in energy than M+H₂0.

Very little gas phase work on reactions of water and methanol with neutral atoms using crossed molecular beams has appeared. Due to the absence of long-range electron transfer, the reactions are expected to have small cross sections ($< 10 \text{Å}^2$). This factor, together with the possibility of several competing reaction channels makes it difficult to obtain a full picture of the reaction dynamics. This is because many detection techniques²² do not have comparable sensitivities to all possible channels. However, a number of interesting features have been observed in gas phase reactions to date. Alcaraz and coworkers⁶ investigated the luminescent channels from the laser-induced reactions of Ba(6s6p¹P₁) and Ba (6a5d¹D₂) with H₂O under single collision conditions. Although no chemiluminescence was observed from reaction of Ba(¹D₂), the Ba(¹P₁) reaction led to BaOH($B^2\Sigma^+$, $A^2\Pi$). No BaO($A^1\Sigma^+$ or $A^{\prime 1}\Pi$) was observed, even though these channels are more energetically favorable. The lack of the concerted molecular channel was attributed to a large barrier on the potential energy surface. The spectroscopy of a number of radicals containing alkaline earth metals has been studied by Bernath²³ and coworkers. Electronic excitation of the metal atom led to substantial enhancement of reactivity: most notably, $Sr(^{1}P_{1}) + CH_{3}OH$ led to formation of $SrOCH_{3}$. However, their focus was on the spectroscopy of the products rather than on the reaction dynamics. A Broida type oven was used in these studies under multiple collision conditions, and the reaction mechanisms could not be unambiguously determined.

Although there have been no reports of formation of MO from single collisions of alkaline earth atoms with water or alcohols, such reactions are known to occur for other atomic species. Gole and Pace¹¹ observed BO chemiluminescence from $B + H_2O \rightarrow BO^{\bullet} + H_2$. Ab initio calculations¹² indicated a large potential energy barrier for this reaction which was apparently surmounted by the fast Boltzmann tail of the effusive atomic B beam. Liu and Parson¹³ studied the reactions of Y and Sc with water and alcohols under single collision conditions. Only the MO product was observed; either the hydroxide was not formed or it's LIF spectrum was outside the wavelength range studied.

We have performed crossed molecular beams studies of the reactions of ground state $(6s^2-{}^1S_0)$ and metastable $(6s5d-{}^1D_2)$ barium atoms with water and its alkyl derivatives. Unlike the previous gas phase experiments, products in our study were detected using an electron bombardment ionizer-quadrupole mass spectrometer. This allowed us to detect all reaction channels, including the dominant ground state products. Our work complements the recent chemiluminescence experiments of Alcaraz et al⁶; the combined results allow us to gain considerable insight into the dynamics of the reactions.

By changing the kinetic energy of the atomic and molecular beams, we compared the effect of reactant collision energy to reactant electronic energy. We find that the reactions with water are largely state specific; at all collision energies, ground state Ba(¹S) reacted primarily to form the molecular products BaO + H₂ by a concerted mechanism, whereas metastable Ba(¹D) preferentially formed the radical products BaOH

+ H. We show that this behavior results from the participation of different electronic potential energy surfaces. Reactions with CH₃OH, on the other hand, only results in formation of BaOCH₃ + H, despite a much larger exoergicity for formation of BaOH + CH₃ or BaO + CH₄ (Table II). Dimethyl ether (CH₃OCH₃) was found to be unreactive even at the highest collision energies. These observations suggest that the formation of radicals results from hydrogen atom migration, a process strongly enhanced by electronic excitation of the incident barium atom.

II. EXPERIMENTAL

A continuous, seeded supersonic beam of barium atoms was formed in a molybdenum oven source with separately heated barium reservoir and nozzle. Details of the apparatus^{5a,24} and the atomic beam source²⁵ are available elsewhere. The molecular beam was formed by bubbling helium or hydrogen carrier gas through the liquid sample held in a refrigerated bath and expanding the mixture through a 0.005° or 0.010° diameter nozzle. A dilute mixture (<5%) and high nozzle temperature (200 °C) was necessary to eliminate dimers and larger clusters. The water was distilled; the alcohols, ether, hydroperoxide and $H_2^{18}O$ were obtained from Aldrich Chemical Company. Both beams were differentially pumped and crossed at 90° in a liquid nitrogen cooled chamber maintained at $\sim 1\times10^{-7}$ Torr.

Reaction products were detected by a triply differentially pumped electron impact ionizer-quadrupole mass spectrometer that could be rotated in the plane of the atomic and molecular beams^{5a,24}. An important goal of the experiment was to measure relative yields

of BaOH(m/e=155) and BaO(m/e=154). This is inherently difficult due to the close proximity of the mass peaks and fragmentation of BaOH to BaO+ upon electron Considerable effort was devoted to characterizing the bombardment ionization. fragmentation patterns of the product molecules and the transmission of the products through the quadrupole mass filter. Fortunately, resolving these products is aided by the favorable kinematic relationships which confine BaO and BaOH to a very narrow scattering angle and lead to rather large signal levels. Also, the background count rate in the detector is nearly zero at these heavy masses. Most importantly, the natural isotopic abundance of Ba is favorable; the isotopes with masses 138, 137, and 130-136 are 71.70%, 11.23% and 17.07% naturally abundant, respectively.²⁶ By operating the mass spectrometer at a high resolution setting, it was possible to distinguish between the ¹³⁸BaOH (m/e=155) and ¹³⁸BaO (m/e=154) parent ion peaks with only a minor contribution at m/e=154 from ¹³⁷BaOH. Fragmentation of BaOH to BaO+ upon 200V electron bombardment ionization was a factor to consider in the analysis. fragmentation ratio is difficult to measure with great accuracy since there are few clean sources of BaOH. The reaction of Ba(1D)+ t-butyl hydroperoxide formed both BaOH + t-butoxy radical and BaOt-Butoxide + OH. Due to the similar exoergicities of the reactions (Table III), and the heavier recoil partner for the BaOH product, it was possible to monitor pure BaOH far from the Ba beam outside the angular range for BaOt-Butoxide. This allowed us to measure the fragmentation pattern obtaining a value of $BaOH^{+}/BaO^{+} = 1.0 + /- 0.2$. This ratio compares favorably with that observed from the product of the reaction Ba(¹D) + H₂O, which we estimate to be 90-100% BaOH.

We have no way to measure the dependence of this pattern on internal excitation of BaOH, but since ionization involves removal of the non-bonding radical electron^{26,27}, it is expected to be small. Adequate discrimination between m/e=154 and m/e=155 was checked by monitoring the BaO product (m/e=154) known to result from the Ba+O₂ reaction^{29,30} and increasing the resolution to extinguish signal observed at m/e=155. It has been found in other work that fragmentation of BaO to Ba⁺ is relatively insensitive to internal excitation.^{5a,31} Product time of flight distributions were obtained using the cross-correlation method, as described previously^{5a,32}.

The metastable barium atomic beam was formed by optically pumping the Ba(¹P¹S) resonance upstream of the interaction region using a Coherent 699-21 ring dye laser operating single frequency at 553nm with Rhodamine 560 dye, as described previously. ^{5a} Based on measurements described elsewhere ^{5a} and using the published branching ratios ³³ for decay of Ba(¹P), the ¹³⁸Ba in the beam consisted of 45% Ba(¹D), 15% Ba(³D) and 40% Ba(¹S), with all other isotopes in the ground (¹S) state. Product angular and velocity distributions were obtained by running parallel experiments at various detector angles with and without the laser. We have not attempted to study the reactivity of Ba(¹P) with water or its alkyl derivatives.

III. RESULTS

A. Reaction of $Ba(^1S) + H_2O$ and D_2O

The reactive and nonreactive scattering of $Ba(^{1}S_{0}) + H_{2}O$ and $D_{2}O$ was studied at several collision energies in the range 12-30 kcal/mole. The Newton diagram for Ba

+ H₂O at a nominal collision energy of 12kcal/mole is shown in Fig. 1. This mean collision energy is just below the 13 kcal/mole endoergicity for Ba(1 S) + H₂O \rightarrow BaOH + H. Assuming that supersonic expansion of 5% H₂O in He at 200°C leads to no vibrational cooling, <2% of the H₂O molecules will have a quantum of symmetric bending excitation (1594cm $^{-1}$). 16 At $E_{coll} = 12$ kcal/mole, only these molecules and the fast edge of each supersonic beam could react to form BaOH + H. However, our experiments indicate that the dominant reaction is actually $Ba(^{1}S) + H_{2}O \rightarrow BaO + H_{2}$. A number of checks were conducted to prove that the BaO signal was in fact due to bimolecular collisions between Ba + H₂O. First, the BaO product angular distribution peaked at the center of mass laboratory angle (20°) consistent with the known masses and velocities of the Ba + H₂O colliding pair. Contributions from reactions of possible contaminants, such as O₂ or (H₂O)_n were ruled out since the center-of-mass angle associated with reaction of these species with Ba is considerably larger. A major concern arose from our observation of trace quantities of BaO in the Ba beam itself. Since the vapor pressure of BaO is extremely low34 at these temperatures, it must arise from reaction of Ba atoms in the beam with residual gasses (O2, H2O) present in the differential pumping region. This BaO impurity remains primarily within the atomic beam. However, we could also detect some BaO outside the beam due to elastic and inelastic collisions between the impurity and water in the molecular beam. In order to eliminate this undesirable signal, a series of experiments were conducted using H₂¹⁸O in place of regular water. Figure 2 shows the Ba¹⁸O time of flight spectra (m/e=156) from reaction of ground state Ba with H₂¹⁸O. The quadrupole mass spectrometer was operated

at unit mass resolution—as expected, no signal could be observed at m/e=156 (Ba¹⁸O) under the same conditions when ordinary water was used in the molecular beam. At this collision energy, the BaO signal level was weak compared to the Ba signal from nonreactive scattering. Figure 3 shows the time of flight spectra recorded at Ba⁺ (m/e=138) under the same conditions. The observed signal primarily resulted from nonreactive elastic and inelastic collisions. The maximum velocity for this nonreactively scattered Ba is shown as a dotted line in the Newton diagram (Fig. 1)—the forward and backward components of the Newton circle can be clearly seen in the Ba time-of-flight spectra (Fig. 3) at lab angles smaller than 30 degrees.

Figure 4 shows the best fit translational energy distribution (P(E)) and center of mass angular distribution (T(θ)) used in the simulation of the data in Figs. 2 and 3 (solid line). A considerable amount of wide angle nonreactive Ba⁺ signal was observed at this collision energy. In figure 3, the minor component (dotted line) in the Ba signal is the contribution from fragmentation of BaO product to Ba⁺ in the electron bombardment ionizer. This contribution was calculated using the best fit to the BaO data and a fragmentation ratio of BaO⁺: Ba⁺ = 1.0. Based on the relative Ba and BaO signal level, we estimate that the cross section for the reaction Ba(1 S) + H₂O \rightarrow BaO + H₂ is < 2Å² at 12 kcal/mole.

Mass scans of the product from the reaction of ground state Ba indicate that formation of BaO + H_2 dominates at all collision energies from reaction of Ba(1 S) + H_2 O. Raising the collision energy above the 13 kcal/mole endoergicity for formation of BaOH + H opened up this channel, but the observed signal level remained substantially

smaller than from BaO + H₂ even at 30 kcal/mole, the highest collision energy studied. Figure 5 shows the angular distributions for the BaO and BaOH from reaction of Ba(1S) + H₂O at a collision energy of 23kcal/mole. Due to the reaction endoergicity and light H-atom recoil partner, the Newton circle for the BaOH product is very small-- the product is constrained to appear near the center-of-mass angle and a substantial fraction of all reaction products enter the detector. In the case of the dominant BaO + H₂ channel however, a smaller fraction of the BaO is collected at a given detector angle since the products are scattered over a much larger Newton sphere. By monitoring both channels at the CM angle (25°), our detection sensitivity is strongly biased towards the BaOH + H channel. This differing kinematic relationship and the occurrence of both channels is clearly illustrated (Fig. 5). At a collision energy of 23 kcal/mole, the width of the laboratory angular distribution observed at BaO⁺ is more than 20 degrees, whereas that for BaOH⁺ is ~10 degrees. Time-of-flight data for the ground state Ba reaction products are shown in figures 6 and 7; the optimized P(E) and T(θ) used to fit the data are shown in figure 8. The contribution to the BaO angular distribution and time-offlights from fragmentation of BaOH to BaO+ in the ionizer is shown in dotted lines in Fig. 7. The laboratory angular distribution for both channels peaked at the center of mass angle (Fig. 5). Based on the fit to the BaO time of flight profiles, (which are considerably broader than those for BaOH) we conclude that there is a substantial release of translational energy in the BaO + H₂ products (Fig. 8). The best fit to the BaO lab angular distribution led to a CM angular distribution which was approximately isotropic. It should be pointed out that due to the kinematic limitation, we would not be able to resolve a slight forward or backscattering. It is even more difficult to make conclusions about the BaOH angular and velocity distribution since the data is more compressed due the light H atom recoil partner. However, our data does indicate that the CM angular distribution cannot be strongly forward or backscattered.

Qualitatively similar behavior was observed when D_2O was used. Again, the BaO + D_2 channel dominated, but BaOD + D was clearly seen at collision energies above the reaction endoergicity. Even at the highest collision energy (30 kcal/mole), nearly that of the Ba(1D) excitation energy, formation of BaO + H_2 (or D_2) remained the dominant channel.

B. Reaction of Ba(1D) with H₂O and D₂O

Excitation of the incident Ba atom to the metastable ¹D state led to a dramatic increase in the observed signal level at all collision energies studied. Whereas the dominant product in the ground state reaction was BaO + H₂ (or D₂), reaction of Ba(¹D) leads primarily, and possibly exclusively, to formation of BaOH + H (or BaOD + D). Figure 9 shows the time of flight spectra obtained at m/e=155 (BaOH). At a collision energy of 12 kcal/mole, the BaOH signal level at the center-of-mass angle (17.5°) from Ba(¹D) was ~20X greater than BaO signal from Ba(¹S). The BaOH laboratory angular distribution and product translational energy distribution are shown in Fig. 10. The large increase in reactive scattering signal can be seen in the Ba time-of-flight data (Fig 11). The dominant contribution (middle peak) to the Ba signal is now from fragmentation of BaOH reaction product, rather than from nonreactive scattering. The Ba time-of-flight

data for collisions of ground and electronically excited Ba are plotted on absolute scales in figure 12. As already mentioned, the large middle component seen with the laser on results from fragmentation of BaOH. The fast component from wide angle nonreactive scattering also increased with electronic excitation. This component likely arises from quenching of electronic excitation.

The dramatic shift in chemical identity of the reaction product upon electronic excitation of Ba is best demonstrated in the product mass spectra. Figure 13 shows the mass spectrum of the products from the reaction of Ba(1S) + D₂0 obtained with the detector at the center-of-mass angle. The collision energy was 23 kcal/mole and the detector was operating at unit mass resolution. A Newton diagram showing the expected angular range for the chemical products is also shown. As before, due to the kinematics, we are most sensitive to the BaOD channel when the detector is at the center of mass angle. Although the observed BaOD intensity from the Ba(1S) reaction is approx. 1/2 that of BaO (Fig. 13, dotted line), the true branching ratio after accounting for this Jacobian factor and the fragmentation of BaOH to BaO+ in the ionization process, is ~ 1:9 BaOD:BaO for Ba(1S) at 23kcal/mole. However, upon electronic excitation of the incident Ba atom to the ¹D state, a large shift in product branching ratio (Fig. 13, solid line) is observed. At 23 kcal/mole, the BaOD signal from the Ba(1D) reaction was approximately 6 times more intense at the center-of-mass angle than the BaO signal from the ground state reaction. With the mass spectrometer tuned to m/e=154 (BaO⁺), the product velocity and angular distribution was found to be almost the same as that observed at BaOD+ and was attributed to BaO+ daughter products from fragmentation of BaOD in the electron bombardment ionizer detector.

From the Newton diagram, the maximum predicted angular range of BaO from $Ba(^1S \text{ or }^1D) + D_2O \rightarrow BaO + D_2$ is substantially greater than from $Ba(^1D) + D_2O \rightarrow BaOD + H$ due to the greater reaction exoergicity and heavier recoil partner (D_2 instead of D). Yet, no BaO could be observed from the $Ba(^1D)$ reaction outside the range where BaOD was seen. Based on this, we conclude that BaOD results from 90-100% of all reactive encounters between Ba (1D) and D_2O . Comparison of signal levels using H_2O and D_2O under similar beam conditions indicated that D_2O is 2.0 ± 0.4 times less reactive than H_2O .

C. Reactions of Ba(1S), Ba(1D) with CH₃OH and CH₃OD.

Three chemically distinct reaction channels, producing BaOH, BaOCH₃, and BaO are thermodynamically possible^{21,23} from reaction of ground state Ba atoms with methanol as shown in Table II. Electronic excitation opens the possibility for formation of electronically excited states of the product molecules²³. Product angular distributions for reactions of Ba with methanol, shown in Fig. 14, indicate that formation of BaOCH₃ + H dominates in reactions of ground state and electronically excited Ba. As with water, strong signal enhancement was seen upon electronic excitation, with the most dramatic increase (20X) at the lowest collision energy studied (15 kcal/mole). At this collision energy, the ground state reaction was only barely detectible. The expected laboratory angular range for the BaOH product is substantially larger than for BaOCH₃ due to its heavier recoil partner (CH₃ instead of H). However, we were unable to detect BaOH⁺

or BaOH daughter products (BaO⁺, Ba⁺, OH⁺) at angles outside of that seen for formation of BaOCH₃ + H, even upon electronic excitation. The observed BaO⁺ and Ba⁺ signal resulted from fragmentation of BaOCH₃ in the electron bombardment ionizer. We conclude that the yield of BaOCH₃ + H is 90-100% from ground state and electronically excited Ba, in spite of the fact that Δ H is more favorable for BaOH + CH₃ production, and formation of BaO + CH₄ is highly exoergic even from the ground state (Table II).

Even at the lowest collision energy studied (15 kcal/mole), ground state Ba atoms were found to react weakly, but again, only forming BaOCH₃ + H (Fig. 14, Top Figure). Spurred by the observation of Ba(lS) + H₂O → BaO + H₂, a search was undertaken for BaO + CH₄, but we failed to see any evidence for this channel from ground state or electronically excited barium up to 30 kcal/mole collision energy. Owing to the large reaction exoergicity and heavier CH₄ recoil partner, some BaO should have been observed outside of the angular range for BaOCH₃ if its formation accounted for more than 10% of the total reaction cross section. Angular distributions for BaOCH₃ from Ba + CH₃OH and Ba + CH₃OD are shown in figure 15. A comparison of Figs. 14 and 15 indicates that the reactivity of Ba(lS) does increase with increasing the collision energy from 15 to 30 kcal/mole. However, electronic excitation is far more effective in enhancing the chemical reaction at any collision energy. A similar isotope effect was observed in the methanol reactions with CH₃OH having a reaction cross section approximately twice that of CH₃OD at the same collision energy

D. Collisions of Ba(1S), (1D) with CH₃OCH₃.

No chemical reaction was observed in collisions of ground state or electronically excited Ba atoms with dimethyl ether at collision energies up to 25 kcal/mole despite long averaging for BaO⁺, BaOH⁺, and BaOCH₃⁺. The observed Ba⁺ signal was attributed to elastic and nonreactive inelastic collisions.

E. Collisions with Allyl Alcohol and t-Butyl Hydroperoxide

In an attempt to observe formation of BaOH from the single collision of Ba + an alcohol, allyl alcohol (CH₂=CH-CH₂-OH) appeared promising. As will be discussed below, the electronic structure of allyl alcohol is very different from that of methanol because of the C=C double bond. Moreover, due to the resonance stability of the allyl radical, formation of BaOH + CH₂=CH-CH₂ is far more exoergic ($\Delta H \sim -28.5$ kcal/mole) than is BaOCH₂-CH=CH₂ + H (Δ H ~ 0 kcal/mole).²¹ Indeed, collisions of Ba(1S,1D) with allyl alcohol resulted in formation of Ba-O-CH2-CH=CH2 + H as well as BaOH + CH_2 =CH- CH_2 . Time-of-flight data for the products from the Ba(1D) reaction are shown in Figs. 16 and 17; the best fit P(E) and T(θ) are shown in Fig. 18. Whereas the Ba-O-CH₂-CH=CH₂ signal is localized near the center of mass angel, (30 degrees) the BaOH product is scattered over a much larger angular range due to its heavier recoil partner (CH₂=CH-CH₂ instead of H). This kinematic relationship is better for the BaOH + CH₂=CH-CH₂ channel than for any of the other channels observed in this study. We find that although the BaOH angular distribution is broad, it is definitely forward peaking, indicating that the reaction occurs on a timescale shorter than ~1psec.

Both channels can be monitored simultaneously at BaO⁺, since both barium containing products fragment in the electron bombardment ionizer. Collisions between Ba and tert-butyl hydroperoxide (t-BuOOH) led to formation of BaO-t-Butoxide + OH and BaOH + t-Butoxide Radical with an approximately 1:1 branching ratio (not shown).

IV. DISCUSSION

A. Reactions with H_2O .

The most striking observation in the Ba + H₂O study was the strong dependence of product branching ratio on initial electronic state of the reactant Ba atom. At collision energies below the 13 kcal/mole endoergicity for Ba(1 S)+ H₂O \rightarrow BaOH + H, the ground state reaction led to formation of BaO+H₂. This channel was very weak at 12 kcal/mole, but increased with increasing collision energy. Although the BaOH + H channel did appear at collision energies above its endoergicity, it remained much weaker than the BaO + H₂ channel even at 30 kcal/mole, a collision energy approaching the Ba(1 D) excitation energy. Electronically excited Ba(1 D), on the other hand, reacted with water to form BaOH+H with a branching ratio of 0.90 - 1.0 at all collision energies.

There are two explanations for the observed product specificity. One possibility is that $Ba(^{1}D) + H_{2}O$ is correlated to BaOH + H through an excited state surface, whereas ground state $Ba(^{1}S) + H_{2}O$ correlates to $BaO + H_{2}$. Alternatively, the effect might be due to energetics alone: electronic excitation may simply drive the endoergic channel. In this case, the collision of $Ba(^{1}D)$ might be viewed as quenching of electronic excitation during the encounter of the reagents with the vibrationally hot

intermediate decaying to form BaOH + H on the ground electronic surface. The very large excess energy above threshold provided by the initial electronic excitation would strongly favor simple unimolecular decomposition from a reaction intermediate leading to BaOH + H rather than formation of BaO + H_2 through a concerted mechanism. However, we believe that this possibility is unlikely in light of the experimental observations . Although BaOH + H was observed from ground state Ba + H_2O at collision energies above its endoergicity, the BaO + H_2 cross section increased and remained dominant with increasing collision energy. If the lack of BaOH + H from ground state reaction was simply due to energetics alone, the large increase in relative translational energy should have led to a shift toward the radical channel, which, although endoergic, should dominate at high energy since it only involves simple H-atom elimination. Since the BaOH + H channel remained weaker than BaO + H_2 even at collision energy near the Ba(1D) excitation energy, energetic arguments alone cannot explain the dominance of BaOH upon electronic excitation.

We now return to the possibility that differing intrinsic features of the ground and excited state surfaces lead to strong preference for different chemical products. Two possible mechanisms can be envisioned for the reaction $Ba(^1S) + H_2O \rightarrow BaO + H_2$. The reaction could be initiated by insertion of $Ba(^1S)$ into one of the OH bonds in H_2O . Although HBaOH is expected to be linear or slightly bent, a very low energy bending potential has been calculated for the analogous molecules involving lighter alkaline earth atoms. ^{7,9} This low energy bending mode could facilitate formation of BaO + H_2 via a four center transition state:

We expect substantial potential energy barriers for this reaction mechanism. The electron configuration for the ground state Ba atom is $6s^2$ - the initial repulsion between this "closed shell" atom and water, as well as the need for sp or sd hybridization for formation of two bonds in the insertion intermediate will likely lead to a substantial potential energy barrier in the entrance channel. Additionally, a second potential energy barrier at the 4-center transition state is expected for H_2 elimination from HBaOH. Simple rupture of the very weak Ba-H bond in H-Ba-OH, on the other hand, should readily lead to BaOH + H with no barrier in excess of the Ba-H bond energy. For these reasons, we believe that an insertion mechanism will lead to BaOH + H, rather than BaO + H_2 . Since BaOH + H was not the dominant ground state product, insertion is not an important mechanism in the ground state Ba reaction.

Our experimental observations, especially the large exit potential energy barrier manifested in a high translational energy release, indicates that a concerted mechanism is involved in the reaction $Ba(^1S) + H_2O \rightarrow BaO + H_2$ with abstraction of the central O atom from H_2O and simultaneous formation of molecular H_2 :

It is well known that concerted reactions are controlled by symmetry constraints.³⁵ We first consider coplanar trajectories in C_{2v} and C_s symmetry. Approach of Ba(1 S) toward H_2O along the C_{2v} (C_s) axis generates a surface of 1A_1 ($^1A'$) symmetry. Similarly, the asymptotic molecular product states with atoms in the same geometry as the reactants generate surfaces of 1A_1 ($^1A'$) symmetry. However, these states do not correlate diabatically, since the ground state of BaO is best represented as $Ba^{2+}O^{2-36}$; the ground state of the product correlates to very high-lying charge transfer states of the reactants. The neutral reactant surface correlates to high-lying non-ionic product states. The avoided crossing of these two 1A_1 ($^1A'$) surfaces generates a substantial potential energy barrier for the concerted reaction—the presence of this barrier leads to the small reaction cross section and rather large translational energy release observed for stable closed shell $BaO + H_2$ products.

In all cases, successful formation of BaO + H₂ requires small impact parameter collisions. This is due to: 1) the very small Ba-O approach distance required for transfer of the first valence electron from ground state Ba to water; and 2) the need to overcome the mechanical barrier associated with sufficient symmetric bending of the water molecule to facilitate concerted formation of BaO + H₂. The covalent-ionic curve crossing radius (R_x) is often applied to reactions of alkali and alkaline earth atoms involving long-range electron transfer. ^{5a,18-20} Although the relation is less applicable in the present case because of the much shorter range for electron transfer due to the negative electron affinity of water, the general conclusions are useful in gaining insight

into the reaction. In units of Angstroms, the crossing point of the ionic and covalent states is given by:

$$R_{x} = \frac{14 \cdot 4}{IP_{Ba} - EA_{M}} (A) ,$$

where IP_{Ba} is the ionization potential of Ba, which is 5.2eV for Ba(^{1}S) and 4.0eV for Ba(^{1}D) 18,19 . EA_M is the 'effective' electron affinity of the molecule (eV), often taken to be somewhere between the vertical and adiabatic electron affinities. As we will show in detail later (section D), all available data indicates that the adiabatic EA_{H20} \leq -2eV³⁷; the vertical EA is near -6eV.³⁸ For Ba(^{1}S), electron transfer requires R_{coll} \leq 2Å. The molecular orbitals for H₂O are: 16

$$(1a_1)^2(2a_1)^2(1b_2)^2(3a_1)^2(1b_1)^2; {}^1A_1.$$

The lowest unoccupied orbital is 4a₁, which is antibonding with respect to the two O-H bonds and bonding with respect to H-H. Only small impact parameter collisions will lead to the first electron transfer into the 4a₁ orbital, at a Ba-O distance of approximately 2Å. This is essentially the bond length of BaO,³⁶ and the second electron transfer required for formation of ground state Ba²⁺O²⁻ will occur readily. Electron transfer into the 4a₁ orbital at such close range with the Ba atom near the symmetry axis

of H_2O will favor symmetric bending of the water molecule and formation of $BaO + H_2$. The need for a small impact parameter collision is consistent with the rather small reaction cross section seen for the ground state reaction.

It is expected that approach of Ba towards the oxygen atom in the plane of the water molecule near the $C_{2\nu}$ axis will most effectively lead to BaO + H_2 since this geometry will enhance symmetric bending. Since the water molecules are randomly oriented, however, most of the collisions involve attack of the Ba atom along trajectories that are not in the plane of the water molecule. In this case, the symmetry is C_1 and all geometries lead to surfaces of A symmetry. However, a substantial barrier is still expected for the same reason as in C_8 and $C_{2\nu}$ symmetries. Of course, small impact parameter collisions will still be required.

Although the light H₂ recoil partner results in compression of the BaO angular distribution, it is either forward-backward symmetric or nearly so. Such forward-backward symmetry is often taken to be evidence for the existence of a collision intermediate with a lifetime exceeding one rotational period (1 psec)^{5a,39-41}. In most cases, such intermediates are associated with deep chemical wells on the potential energy surface. In the present case, the Ba-H₂O interaction is not much stronger than a typical dispersion interaction with a binding energy not greater than 10 kcal/mole⁷. It might thus appear surprising that a large fraction of the colliding partners interact long enough to undergo at least one half of a rotation before product formation. However, a concerted three-center elimination of H₂ requires sequential two electron transfer with considerable

deformation of water. This necessitates a closely coupled collision with the Ba reaching the repulsive inner wall of the Ba-H₂O interaction potential before the initiation of electron transfer. Although the Ba-H₂O interaction is not very strong, there likely exists entrance and exit potential energy barriers which lead to an effectively deeper well. Also, in the absence of significant steric effects, the CM angular distribution will be broad since non-coplanar trajectories are statistically dominant.

Symmetry arguments may be made for interaction of $Ba(^1D) + H_2O$; however, the situation is rather more complicated. In C_{2v} , the following surfaces are generated for the reacting pair: 1A_1 , 1A_1 , 1A_2 , 1B_1 , 1B_2 . According to Douglas et al⁸, the lowest lying states are likely to be 1A_1 and 1A_2 , the former state having the same symmetry as the ground state molecular products $BaO + H_2$. But again, a substantial barrier is expected to be generated by the crossing of the excited state reactant and ground (or excited) state product surfaces; the existence of a barrier for formation of $BaO + H_2$ (or $BaO^* + H_2$) on the excited surface is consistent with the lack of enhancement of BaO formation upon electronic excitation in both our experiment and in that of Alcaraz, et. al.⁶

We now turn to the mechanisms expected for formation of the radical products

BaOH + H. Our observation that this channel is promoted by electronic excitation

rather than collision energy is good evidence that the reaction occurs much more readily

on an excited state surface. This possibility was first suggested by Douglas and

Margrave⁷ in discussion of the matrix reactions of alkaline earth atoms with water. In

their experiment, irradiation of the Ba-OH₂ adduct, resulting from co-condensation of Ba—

+ H_2O in a matrix, near the Ba atomic resonance facilitated "insertion" of the metal atom into the OH bond with subsequent formation of BaOH. The question naturally arises as to whether the gas phase reaction of electronically excited Ba is mediated by an insertion mechanism, and if so what is the nature of the motion of the atoms during the process? The results of the alcohol experiments are very helpful in our understanding of this radical channel. We turn to those results before returning to a discussion of the $Ba(^1D) + H_2O$ reaction mechanism.

B. Reaction with CH₃OH

The reaction of ground state and electronically excited Ba with CH₃OH shows remarkable preference for formation of BaOCH₃ + H. The other product channels BaOH + CH₃ and BaO + CH₄ either do not exist, or are so weak as to be below the sensitivity of our experiment. We conclude that the hydrogen atom elimination channel accounts for between 90 and 100% of all reactive encounters in the case of ground- and excited state Ba + CH₃OH.

It is worth noting several important differences between H₂O and CH₃OH that pertain to reactions with Ba. The most obvious difference is that formation of BaOH + H is endoergic for ground state Ba, whereas formation of BaOCH₃ is nearly thermoneutral. We did observe formation of BaOCH₃ from reaction of ground state Ba even at the lowest collision energy obtainable in this experiment, although weakly. Our observation that BaO + CH₄ was not detected from the Ba(¹S or ¹D) + methanol

reaction, whereas $BaO + H_2$ did result from collision of $Ba(^1S)$ with water, is not too surprising. Formation of $BaO + CH_4$ would probably have to be of a concerted nature since insertion would likely favor radical production, as outlined earlier for the $Ba + H_2O$ reaction. Such a concerted reaction is less likely in the case for methanol than water since it requires attack by the H atom from -OH on an sp^3 hybridized carbon atom via a three-center transition state:

The potential energy barrier for this process will be very high since at the transition state, the strong O-CH₃ bond has to be nearly broken to facilitate bonding between the tetravalent carbon and the attacking H-atom. However, for trivalent group IIIA elements such as Sc and Y, formation of MO + CH₄ was observed by Liu and Parson¹³ under single collision conditions. They were unable to observe formation of the metal hydroxides-- either they were not formed or their LIF spectrum was outside the wavelength range studied.

Upon electronic excitation, the large increase in formation of BaOCH₃+ H from methanol is analogous to the increase in BaOH + H from water. The most obvious explanation for the formation of BaOCH₃ + H rather than BaOH + CH₃ at all collision energies is that the reaction is dominated by internal migration of the H atom from O to

Ba within the Ba-O bonded intermediate:

$$H_3C$$
 H_3C $\\ O - Ba \rightarrow BaOCH_3 + H$ $\\ H$

We believe that a mechanism involving H-atom migration will involve substantially smaller potential energy barriers than alternative mechanisms leading to HOBaCH₃, ie CH₃ migration or insertion of Ba into the HO-CH₃ bond. This is largely due to the spherical shape of the H atom 1s orbital: good orbital overlap may be maintained with the O and Ba atoms simultaneously during the H-atom migration process. This situation contrasts the directional nature of the sp³-hybridized methyl group which necessitates nearly complete rupture of the O-CH₃ bond prior to formation of HOBaCH₃.

The present reaction is somewhat analogous H₃C-H and H-H bond activation by transition metals such as platinum and palladium.^{42,43} It has been found that the potential energy barrier is very small for insertion of these metals into H₂. However, a substantial barrier exists for formation of H-M-CH₃-- it is much larger for formation of CH₃-M-CH₃. This behavior has been rationalized in terms of the ability for the spherical H-atom to participate in multicenter bonding.^{42,43} Multicenter bonding is less feasible for -CH₃ because the tetrahedral sp³ hybridization leads to highly directional bonding capability. Our observation that dimethyl ether is unreactive under single collision conditions also

supports our hypothesis that H-atom migration is the dominant mechanism in the radical channels.

This work appears to be the first direct evidence of a metal methoxide in the gas phase under conditions known to involve only a single collision. As far as we know, only one previous $M + CH_3OH$ reaction has been studied in rare gas matrices. Codeposition of $Fe + CH_3OH$ led to formation of an $Fe(CH_3OH)$ adduct with the iron atom bonded to the oxygen atom in methanol. Irradiation of the adduct at $400 \text{nm} < \lambda < 500 \text{nm}$ resulted in formation of HFeOCH₃, whereas irradiation at $280 \text{nm} < \lambda < 360 \text{nm}$ led to $HOFeCH_3$. In other words, a higher photon energy led to 'insertion' into the C-O bond. Although the original authors did not draw any conclusions about the reaction mechanism, we suggest that H-atom migration is likely to be the lowest energy process in the iron system. It would not be surprising if this was a general feature in reactions of metal and main group atoms with H-atom containing compounds.

C. The effect of deuterium substitution and collision energy on Ba(1D) reactions.

Comparison of BaOH to BaOD signal levels indicated that $Ba(^1D) + D_2O$ was 2.0 ± 0.4 times less reactive than the H_2O reaction; this ratio was found to remain constant to within the stated uncertainty as the collision energy was varied from 13-30 kcal/mole. The overall reactive cross section also remained constant to within the experimental uncertainties over this range.

The observed isotope ratio observed here is not large by comparison to low temperature experiments where the "semiclassical effect" (i.e. without the occurrence of tunneling) is commonly in the range of 3 to 5.45,46 Experimental observations and theoretical calculations on the Si + H₂O system have indicated that reaction in rare gas matrices occurs with H-atom migration by quantum mechanical tunneling. Co-condensation of Si + H₂O led to rapid formation of HSiOH, whereas visible irradiation of Si + D₂O was required for formation of DSiOD. If the barrier for the rate limiting step in the reaction of Ba (¹D) + water was large, it might be possible to observe a substantially increased isotope effect with decreasing collision energy if tunneling were important. We have found that the isotope effect remains constant over our experimentally obtainable collision energies. Thus, it appears that at these collision energies, quantum mechanical tunneling is not a dominant factor in the deuterium isotope effect that we observe.

D) Product Angular Distributions for the Radical Channels

Although the BaOH and BaOCH₃ angular distributions are rather compressed due to their light H or D atom recoil partner, the best fit of the angular and TOF data was found to employ a forward backward symmetric, isotropic center-of-mass angular distribution (to within the sensitivity of the fit). The usual explanation for this angular distribution is the existence of an intermediate HBaOR species with a lifetime exceeding one rotational period. It should be noted, however, that the symmetric product angular

distribution might also be attributable to the migration process which by its very nature is likely to reflect the symmetry of the water molecule:

Since both H atoms are equally likely to migrate, a forward-backward symmetric angular distribution will result even if HBaOH has a lifetime of less than one rotational period.

E. Comparison with dissociative electron attachment experiments-

Considerable insight into the similarities and differences between H₂O, CH₃OH and CH₂=CH-CH₂OH can be gained by examining results of dissociative low energy electron attachment experiments. Electron attachment to water was studied by Melton³⁸ in the early '70s; the results on methanol and allyl alcohol are due to Illenberger, et. al.⁴⁸ Briefly, the negative ion fragments resulting from dissociation of the temporary compound ion states are detected as a function of incident electron beam energy. Such experiments provide information on the energies of unoccupied molecular orbitals as well

as favorable dissociation channels. As we have already mentioned, long range electron transfer cannot occur for this class of reactions, even with electronically excited Ba. However, it is clear that a transition must occur between the covalent surfaces which characterize the reactants and those of the products which are definitely ionic.

When low energy electrons are passed through water vapor, three rather weak resonances are observed at 6.4, 8.4 and 11.2 eV; all lead to formation of H', OH and O.38 The existence of three peaks has been attributed to three electronic states of H₂O, namely ${}^{2}B_{1}$, ${}^{2}A_{1}$ and ${}^{2}B_{2}^{38,49}$; apparently, all dissociate to yield the three ions. In the case of methanol, again signal was observed at three energies—6.2, 7.8 and 10.2ev, i.e., the energy levels of the unoccupied valence orbitals are comparable for water and methanol. However only CH₃O was produced at 6.2 eV, and CH₃O dominated at 7.8eV; but all three species O', OH', and CH₃O are produced at the highest energy⁴⁷. Allyl alcohol, on the other hand exhibited a much lower energy resonance than water or methanol, at 2eV. Whereas the lowest energy electron attachment to methanol led only to CH₃O, dissociation of the allyl alcohol anion led to both OH and CH₂=CH-CH₂O. It appears that a direct analogy may be drawn between the negative ions resulting from decay of the lowest energy resonances in electron scattering with water, methanol, and allyl alcohol, and the negative ions in the radical products resulting from their reactions with Ba atoms.

The exclusive formation of CH₃O rather than OH from electron attachment to methanol at 6.2eV cannot simply be attributed to thermodynamic effects since the exclusive channel is the most endoergic:⁴⁷

e + CH₃OH
$$\rightarrow$$
 CH₃ + OH, Δ H_o = 2.1 ev,
 \rightarrow CH₄ + O, Δ H_o = 2.4eV,
 \rightarrow CH₃O + H, Δ H_o = 2.9eV,

When several dissociative channels are available, usually the fragment with the highest electron affinity retains the electron.³⁸ This is not true in the present case-- the electron affinity of CH_3O $(1.57\pm0.022eV)^{50}$ is less than that of $OH(1.829\pm0.010eV)^{51}$. This suggests that the electron scattering behavior results from dynamic, rather than thermodynamic considerations. Similar behavior is seen in photodissociation of alcohols. The photodissociation of CH_3OH at 193nm is a direct process preferentially resulting in H atom elimination, whereas formation of vibrationally activated ground state CH_3OH by insertion of $O(^1D)$ into CH_4 primarily led to the thermodynamically most favored products, $OH + CH_3^{52}$.

Based on these observations, it would be tempting to suggest that the lowest lying unoccupied molecular orbital of methanol is primarily of O-H antibonding character-electron attachment or electronic excitation of methanol leads to preferential H atom elimination. Since a switch to an ionic surface inevitably occurs in the Ba + CH₃OH reaction, this might also explain the dominance of BaOCH₃ + H. According to Ab-Initio calculations, ^{53,54} the situation actually somewhat more complex. It appears that the first excited state of methanol is repulsive in both the O-H and C-O coordinates. However, there is a barrier in the adiabatic potential curve for the C-O dissociation coordinate

arising from a sharp Rydberg-valence avoided crossing.⁵³ In contrast, the O-H dissociation coordinate was found to be completely repulsive.

The very light H atom product in the $CH_3O^- + H$ (or $CH_3O + H$) channel leads to an important dynamic effect. For a given recoil energy, the relative velocity of the products ($\approx v_H$) will be five times greater than for $OH^- + CH_3$, (or $OH + CH_3$). Thus, the dissociation dynamics primarily derives from the steeper potential in the O-H coordinate and the much greater relative velocity for $H + CH_3O$. In the $Ba + CH_3OH$ reaction, similar effects, together with a lower potential energy barrier for H-atom migration, lead to preferential formation of $BaOCH_3 + H$ rather than $BaOH + CH_3$.

The reactivity and low energy electron scattering with allyl alcohol contrasts the Ba + H₂O and CH₃OH systems. In this case, the low energy resonance is nearly identical to that in ethylene.⁴⁸ Electron attachment to the low lying π_{CC}^{\bullet} molecular orbital results in rapid randomization of the excitation energy followed by dissociation to OH or OCH₂-CH = CH₂·.⁴⁸ We propose that an analogous situation occurs in the reaction of allyl alcohol with Ba; apparently formation of BaOH + CH₂-CH=CH₂ (favored by thermodynamics) can compete with formation of BaOCH₂-CH=CH₂ + H (favored by the mobility of the H-atom).

E. The role of the excited state in H-atom migration.

As was pointed out by Douglas and coworkers⁷, HBaOH can be treated by analogy with the HAB molecule, with B = OH. We discuss the particular case of water

here: a parallel argument should apply to the reaction of methanol since substitution of H by CH₃ should have little effect on the electronic structure of the Ba core. For HBaOH, the ground state configuration is 'A' (1a')2(6a')2, with an excited state (1a')2(6a')(7a'), also of 1A' symmetry. In the case of the heavier alkaline earth atoms, the ground state 6s-a' orbital of the metal atom probably lies higher than the unoccupied 4a₁-a' orbital of water. The ground state configuration of Ba+ H₂O in C₂ symmetry correlates diabatically with an excited state of HBaOH, whereas the ionic state Ba+H2O-(or possibly Ba2+H2O2-) correlates to the HBaOH ground state. An avoided crossing of these two curves generates a substantial barrier for the ground state insertion process. We observed the onset of Ba(1 S) + H₂O \rightarrow BaOH + H at collision energies above 13 kcal/mole. This might indicate that the barrier on the ground state surface does not lie much above the reaction endoergicity. Alternatively, the ground state reaction Ba(1S) + H₂O → BaOH + H might result from a direct abstraction mechanism, rather than from H-atom migration. In any case, it is clear that the potential energy barrier for formation of BaOH + H is higher than that for formation of BaO + H2, which was observed to occur readily even at the lowest collision energy (12 kcal/mole).

Our observation that electronic excitation is much more effective than collision energy in enhancing the formation of BaOH + H is good evidence that the barrier for formation of H-Ba-OH by H-atom migration is low on the excited state potential energy surface. There has recently been considerable theoretical work on the activation of covalent bonds by transition metals. 42,43 The reaction M + H-R \rightarrow H-M-R (R=alkyl) is

most favorable for those metal atoms with ground state electronic configurations favorable to sd hybridization, since this readily facilitates formation of two covalent bonds. Bond activation is also favored by electronic configurations which minimize long-range electronic repulsion. In the present case, the 6s5d(¹D) configuration has the "singlet diradical" character favorable for formation of H-Ba-OR. This is clearly not the case for Ba 6s²(¹S)-- the paired 6s² electron configuration will have a substantial promotion energy. It will also have a substantially weaker interaction with water due to its closed shell character.

The main difference between Ba + H-OR \rightarrow H-Ba-OR and transition metal reactions such as M + H-R \rightarrow H-M-R (R=H,CH₃) is that the Ba intermediate has substantial ionic character, whereas the transition metal intermediate is primarily covalently bonded. Thus, some additional insight into the reaction dynamics may be gained by considering the factors which favor the transition from the covalent to ionic surface which must ultimately occur for formation of Ba⁺OR⁻+H. Vertical electron transfer to H₂O (especially from Ba(1 S)) is thermodynamically unfavorable, with the lowest vertical electron attachment resonance at 6.4eV.³⁸ Interestingly, the literature reveals one experimental observation³⁷ of H₂O⁻ in ion cyclotron resonance experiments. It has been determined that this molecule, probably best described³⁷ as OH⁻...H is stable and lies about 2.1 eV above H₂O + e⁻. This is below the energy of separated OH⁻+H (3.28eV)²¹. The Ba-water collision timescale is long compared with the vibrational frequency of water. Electron transfer from Ba to H₂O will be most favorable for

collision geometries resulting in asymmetric stretching of the water molecule. For those molecules, the "effective electron affinity" will be near -2.1 eV, and the curve crossing radius for Ba(1 D) is ~2.5Å. Note that the crossing radius based on eq. 1 for Ba(1 S) will still be below 2Å even with this "most favorable" value of -2.1 eV for the EA of H₂O. This difference between the curve crossing radii in these two cases will play a role in determining the identity of the final product. In the case of electronically excited Ba, transfer of the more easily ionized d-electron to the symmetry-allowed orbitals in H₂O will preferentially occur at ~2.5Å in collisions leading to asymmetric stretching of the water molecule. The weakly bound H atom in the newly formed H₂O molecule will readily migrate away from the oxygen before the Ba can approach to within the close range required for second electron transfer that is required for formation of BaO($^{1}\Sigma$) + H₂.

V. CONCLUSIONS

The reaction of Ba with water illustrates that under appropriate conditions, it is indeed possible to control the chemical outcome of a reaction by proper selection of the initial electronic state of a reactant. The observation that reaction of ground state Ba results in BaO + H₂, whereas Ba electronic excitation leads to BaOH + H, is a consequence of the different dynamic and electronic requirements for each channel. Due to the existence of a large mechanical barrier, and the need for close approach of the reactants, the molecular products are favored by the ground state barium atoms. For

only in the ground state can Ba approach to within ~2Å before the first electron transfer. Concerted molecular rearrangement with second electron transfer can occur near the inner turning point of the collision, resulting in production of BaO + H₂. Formation of radical products, on the other hand, results from H- atom migration. Initial excitation of one valence electron to the 5d-orbital strongly promotes this process. Two main factors contribute to this effect. The "singlet diradical" character of the 6s5d configuration is more favorable to formation of two bonds required for H-Ba-OR. Electronic excitation also increases the electron transfer radius. Electron transfer at longer range is favored by asymmetric stretching of the water molecule, leading to H-atom migration and radical formation. This precludes the second electron transfer required for the molecular channel.

The observation of only BaOCH₃ from reaction with methanol, and lack of reaction of dimethyl ether has provided additional evidence for the importance of H-atom migration in the radical channels. The parallels drawn between reactions involving electron transfer, and the products from dissociative electron attachment, suggests that analogies between the processes may provide a solid basis for future predictions of the most probable outcomes of other chemical reactions, which in principal might posses more than one possible ionic product.

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intermediates not connected to lines were not observed in this work or in chemiluminescence experiments described in Ref 6.

Fig20 Same as Fig. 2, but reaction is Ba + CH₃OH.

Table 1. Thermodynamics of the Ba + H_2O System^{6,7,18}

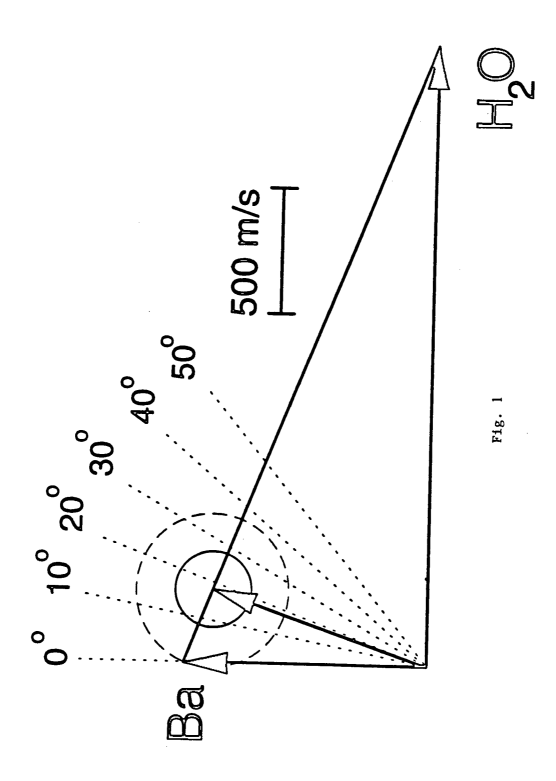
Reaction		$\Delta H(kcal/mole)$
$\frac{1}{\text{Ba}(^{1}\text{S}) + \text{H}_{2}\text{O}}$	→ Ba-OH ₂	-5
	→ HBaOH	-29
	\rightarrow BaOH(X) + H	+13
	\rightarrow BaO(X) + H ₂	-17
$Ba(^{1}D) + H_{2}O$	→ BaOH(X) + H	-20
	\rightarrow BaO(X) + H ₂	-49
	\rightarrow BaO(A,A') + H ₂	≈ 0

Table 2. Thermodynamics of the Ba + CH_3OH System^{6,7,18}
Reaction $\Delta H(kcal/mole)$

		•
Ba(¹ S) + CH ₃ OH	→ Ba-OH(CH ₃)	-5
	→ HBaOCH ₃	-29
	→ HOBaCH ₃	-29
	→ BaOH(X) + CH_3	+13
	\rightarrow BaO(X) + CH ₄	-17
Ba(¹ D) + H ₂ O	\rightarrow BaOH(X) + H	-20
	\rightarrow BaO(X) + H ₂	-49
	\rightarrow BaO(A,A') + H ₂	≈ 0

Table 3. Thermodynamics of Other Systems Studied^{6,7,18}

Reaction		Δ H(kcal/mole)	
$Ba(^{1}S) + CH_{3}OCH_{3} \rightarrow BaOC$	$CH_3(X) + CH_3$	+13	
→ BaO(2	$X) + C_2H_6$	-17	
$Ba(^{1}S) + CH_{2} = CHCH_{2}OH \rightarrow BaOH(X)$		CH ₂ =CHCH ₂ -20	
-	→ BaOCH ₂ CH ₂ CH ₃	+ H -49	
$Ba(^{1}S) + (CH_{3})_{3}COOH \rightarrow Ba(^{2}COOH)$	OH + (CH ₃) ₃ CO		
→ BaO	$(CH_3)_3 + OH$		



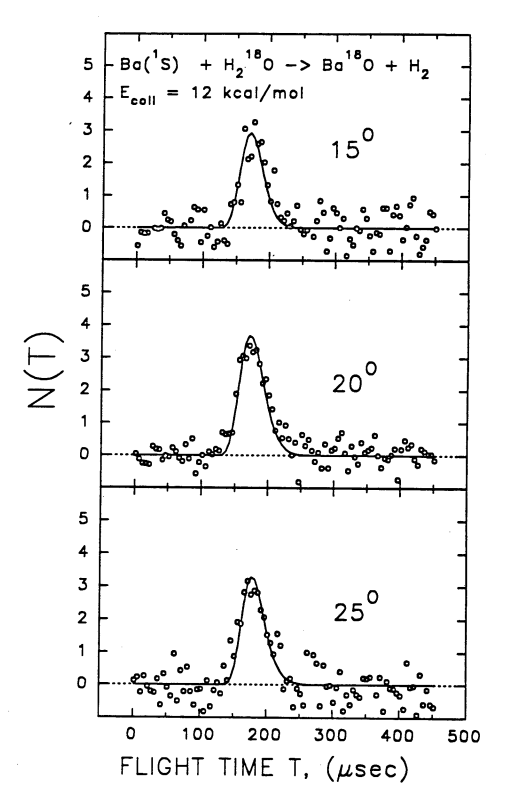


Fig. 2

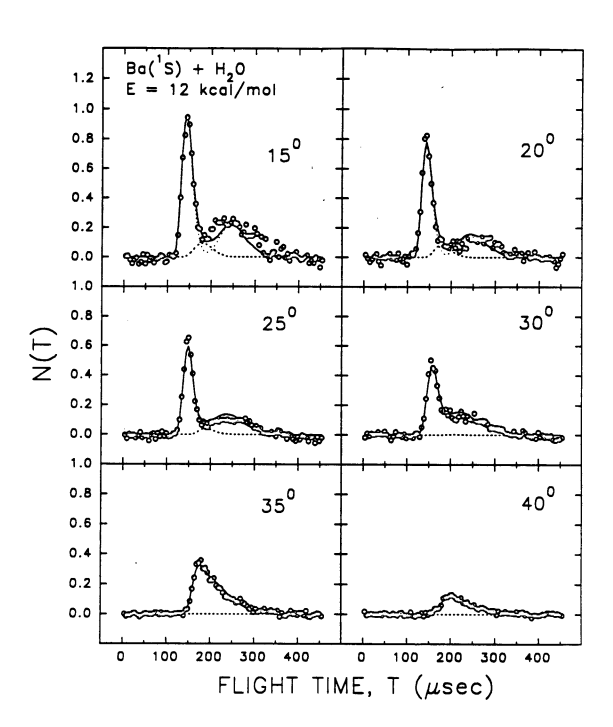
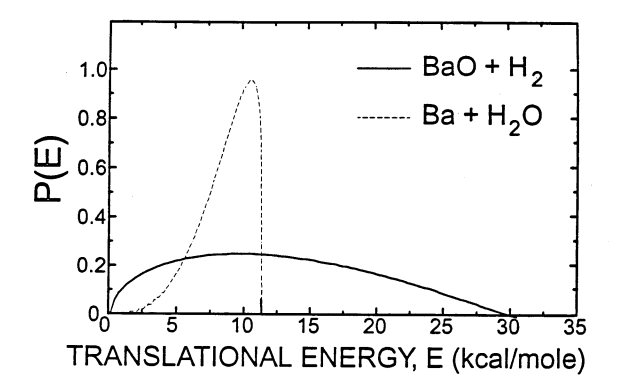


Fig. 3



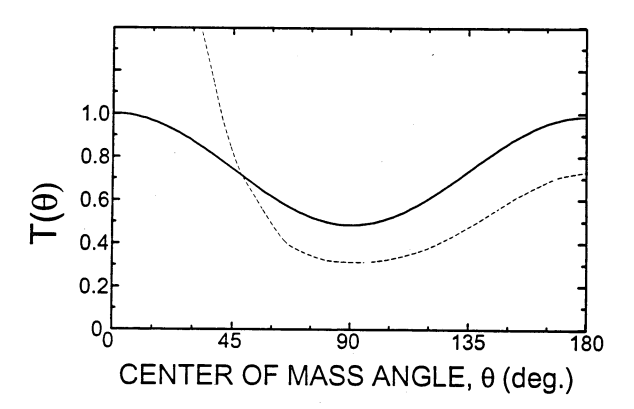


Fig. 4

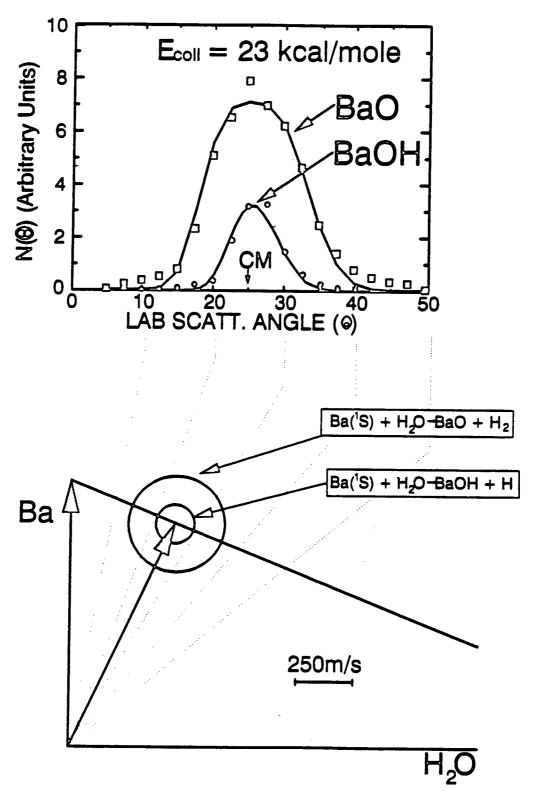
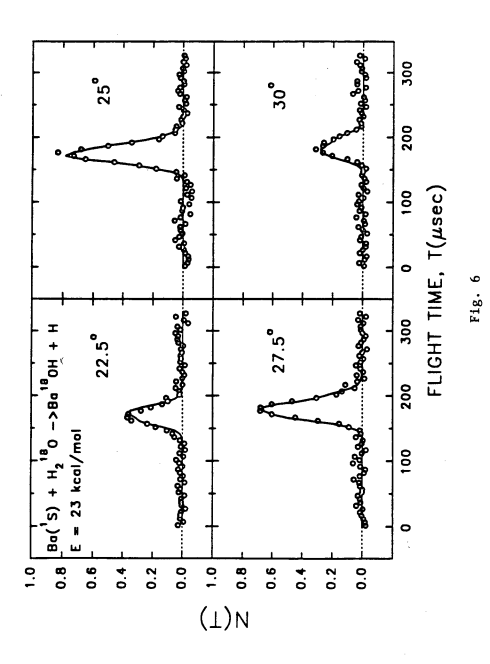


Fig. 5



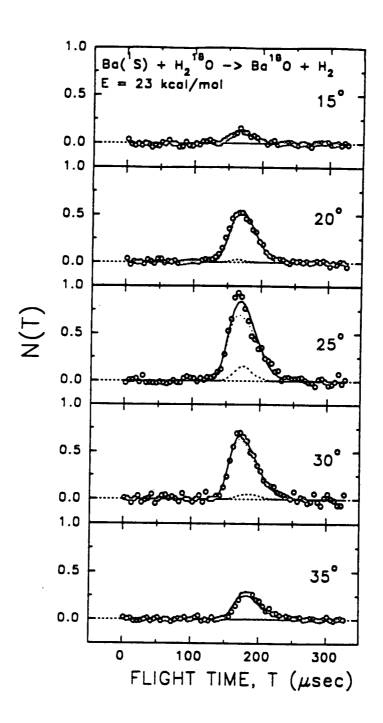


Fig. 7

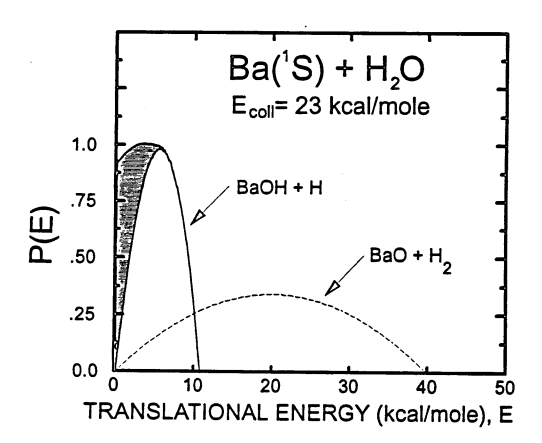


Fig. 8

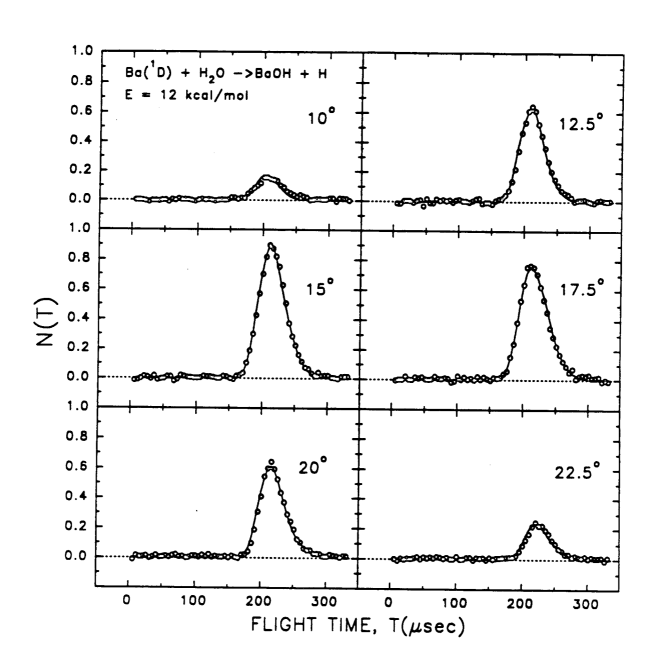


Fig. 9

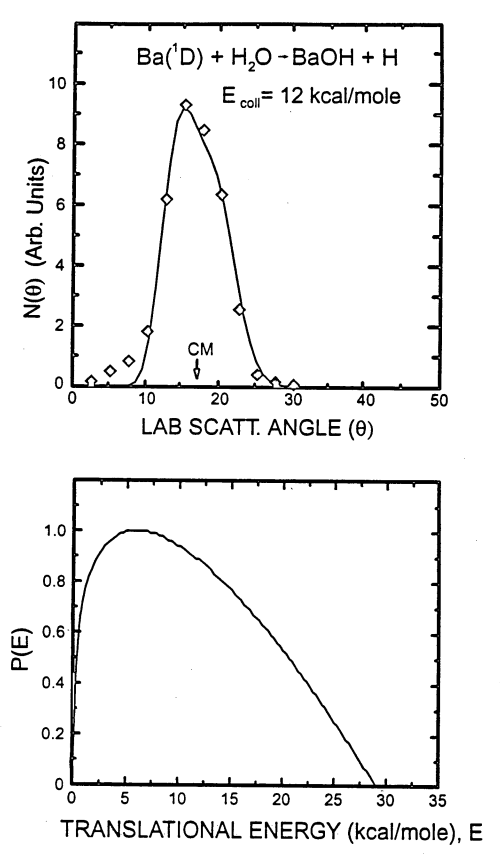


Fig. 10

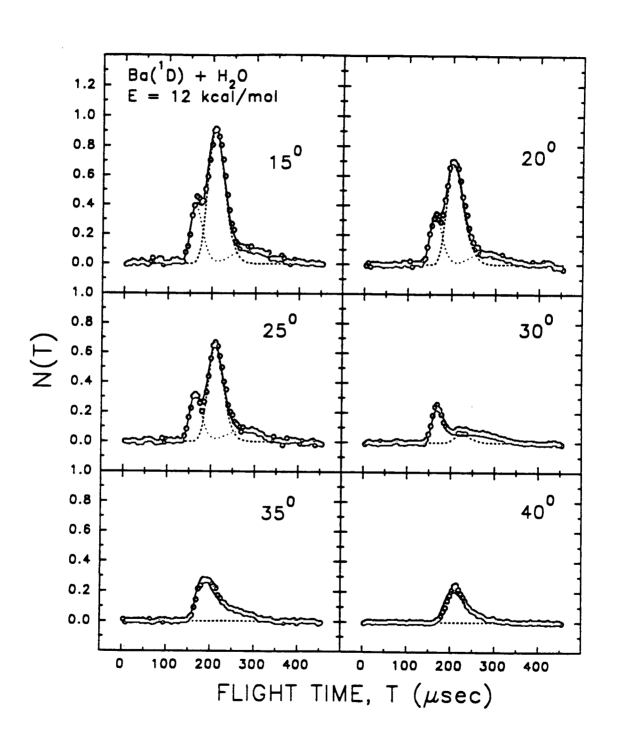


Fig. 11

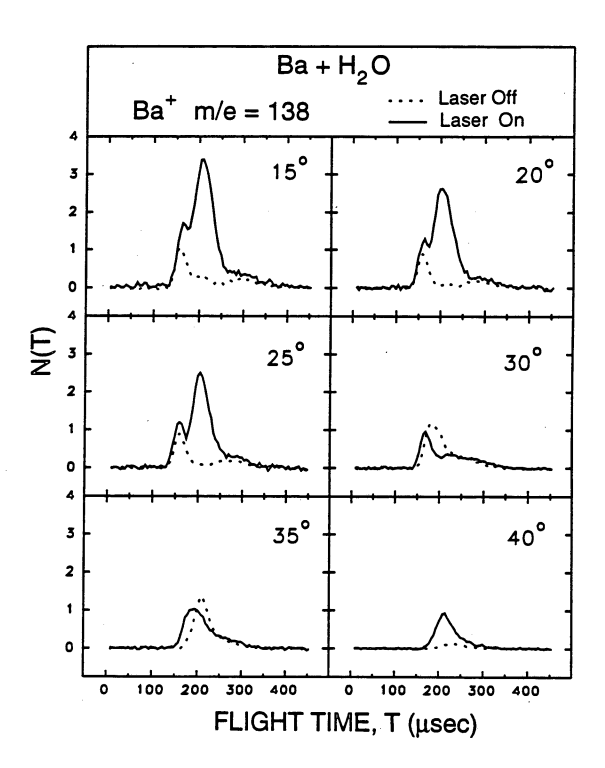


Fig. 12

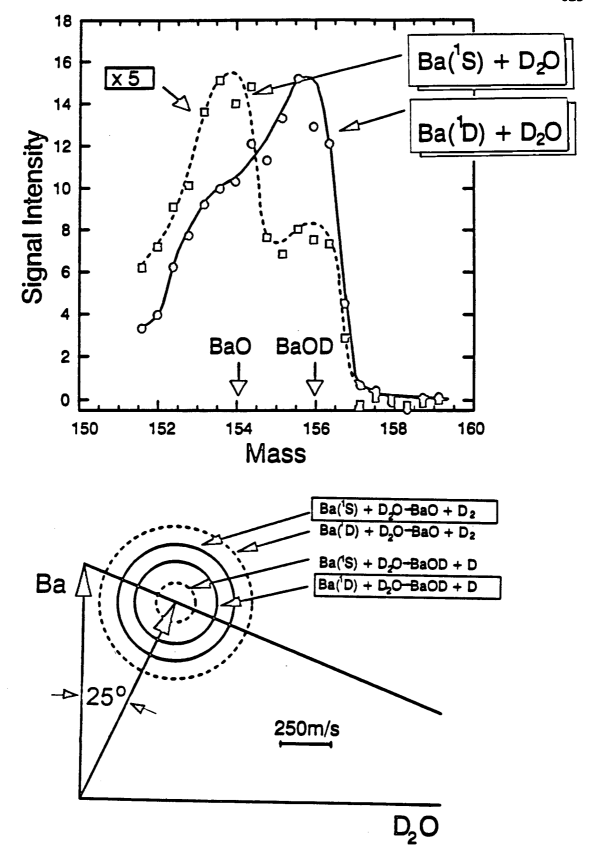


Fig. 13

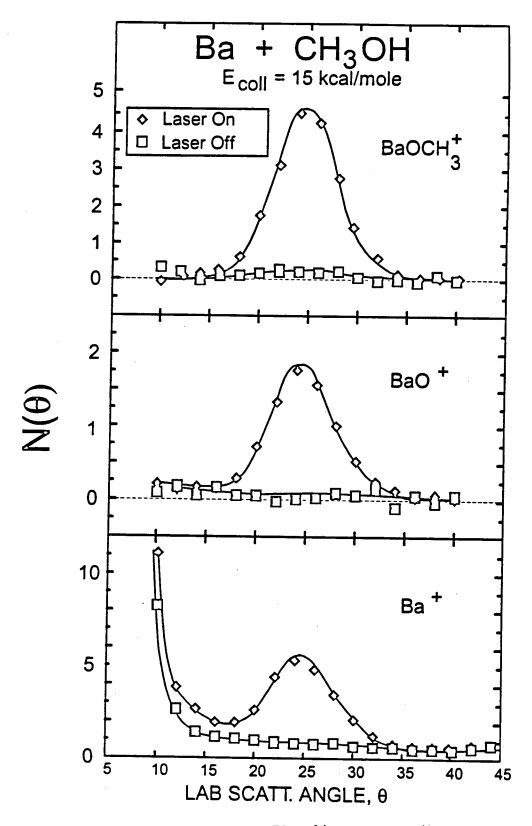


Fig. 14

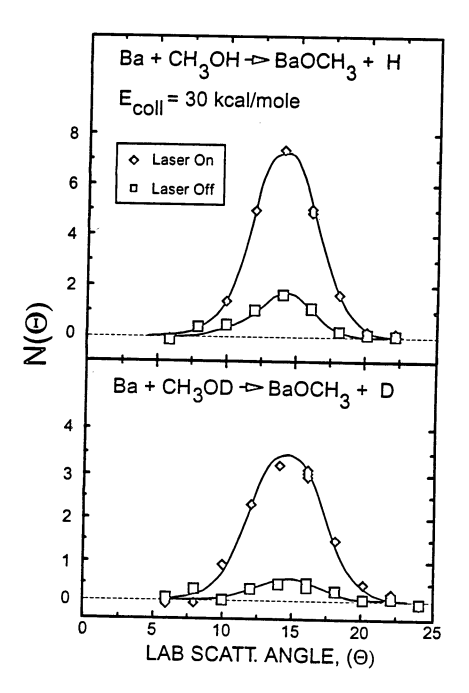


Fig. 15

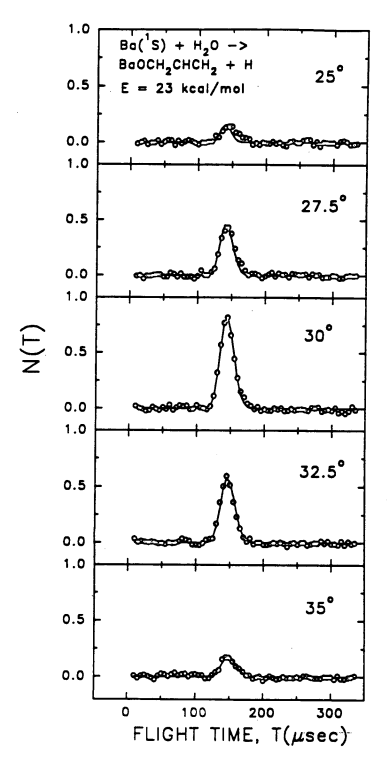


Fig. 16

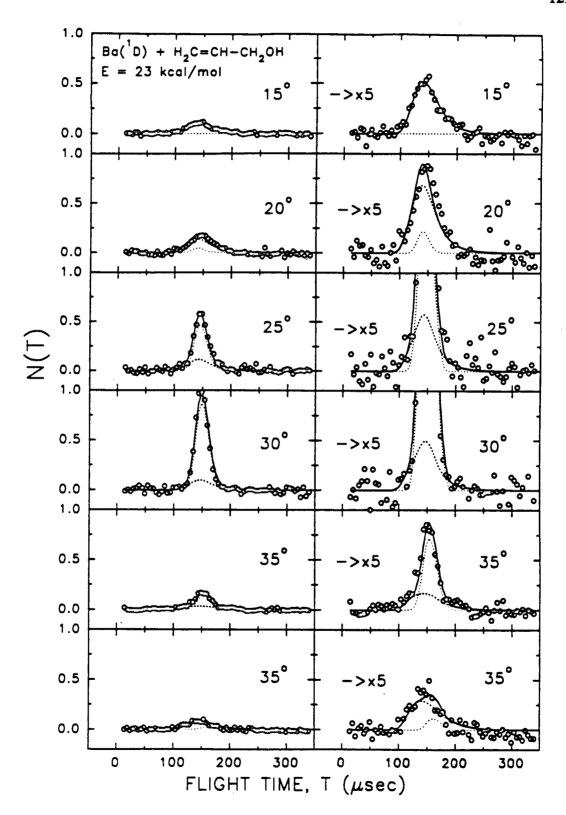
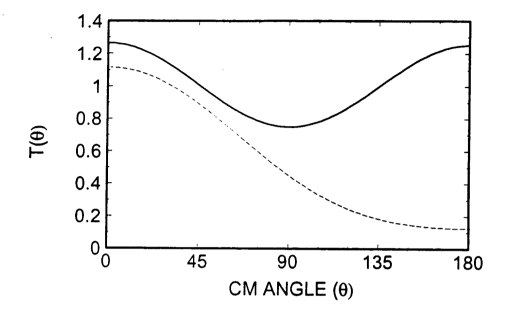


Fig. 17



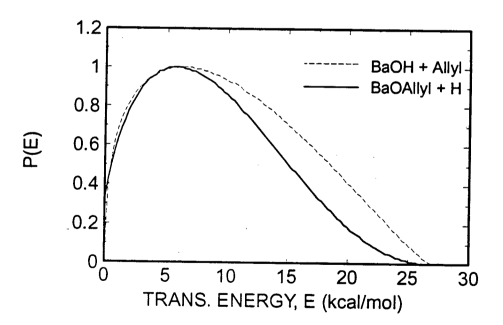
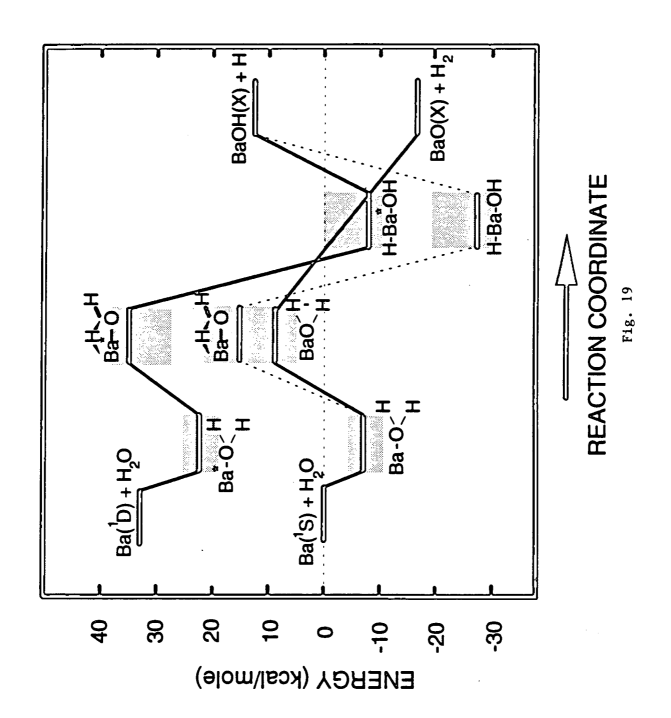
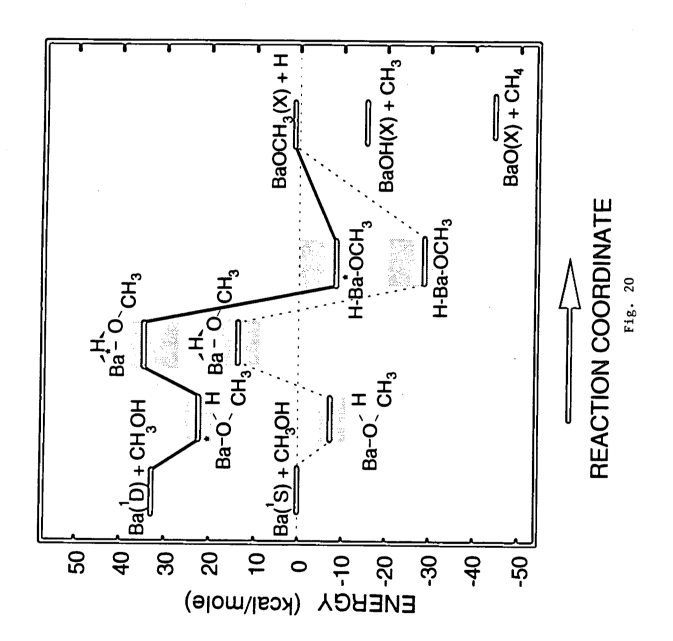


Fig. 18





CHAPTER 3

Reactions of Ba atoms with Triatomic Oxidants: Ba + ClO₂ and O₃

ABSTRACT

This chapter presents the results of a study of the reactions of Ba atoms with symmetrical chlorine dioxide (ClO₂) and ozone (O₃) using the crossed molecular beams method. As was the case in Ba + NO₂ (Chapter 1), chemical products result from direct reaction mechanisms, and by decay of long-lived collision complexes. Single collisions between Ba + ClO₂ led primarily (\sim 90%) to formation of BaO + ClO. The dominant mechanism involved an ionic Ba⁺ClO₂ intermediate that survived longer than one rotational period. Formation of BaCl + O₂ by abstraction of the central chlorine atom of ClO₂ involving a concerted mechanism accounted for \sim 10% of the total reaction cross section. The BaCl angular distribution indicated that the Ba + ClO₂ collision partners survive for at least several rotational periods before concerted decomposition to BaCl + O₂. We observe no evidence for formation of BaO₂ + Cl, the most thermodynamically favorable channel.

Two chemically distinct reaction channels, $BaO + O_2$ and $BaO_2 + O_3$, resulted from $Ba + O_3$. This reaction led to a much larger fraction of forward scattered BaO_3 than was seen in the NO_2 and ClO_2 cases, indicating a greater contribution from direct reactions. This is consistent with the decreased thermodynamic stability of O_3 relative to NO_2 and ClO_2 upon electron attachment to the neutral molecule. This work represents the first direct observation of nascent BaO_2 in the gas phase. Based on the

maximum $BaO_2 + O$ translational energy release, we calculate $D_o(Ba-O_2) = 120 \pm 20$ kcal/mole. This is nearly twice that of comparable alkali superoxides, (e.g. $Cs^+O_2^-$), suggesting that BaO_2 possesses considerable peroxide (doubly ionic) character. Despite the very large reaction exoergicity (~98kcal/mole) for $Ba + O_3 \rightarrow BaO_2 + O$, the reaction results exclusively from decay of long-lived $Ba^+O_3^-$ collision complexes. This is attributed to the existence of a large exit potential energy barrier associated with the electronic rearrangement required for decay of the initially formed singly ionic $Ba^+O_3^-$ to doubly ionic $Ba^2^+O_2^{-2}$. The $Ba + O_2$ reaction is discussed in light of these new results.

I. INTRODUCTION

Walsh's rules³ and molecular orbital theory⁴ allow us to predict trends in the geometries of triatomic AB₂ species as well as changes in bond angles and bond lengths upon electronic excitation or electron attachment. Based on the success of these ideas, it seems natural that systematic trends in the dynamics of reactions initiated by long-range electron transfer are likely to exist in the series of bent triatomic molecules: NO₂, O₃ and ClO₂. With this in mind, we have conducted a systematic study of the reactions of Ba with these oxidants using crossed molecular beams. The NO₂ reaction was discussed in Chapter 1: we now focus on reactions with O₃ and ClO₂. The ultimate goal of our work is to further our understanding of the general principles important in the reactions of divalent atoms with triatomic oxidants.

Several studies on the gas phase reactions of barium atoms with ozone⁵⁻⁸ and

chlorine dioxide⁹ have appeared in the literature over the past twenty years. In all cases, only the chemiluminescent or ionic products were studied although it was known that under single collision conditions, these channels represent no more than 20% of the total reaction cross section.⁶⁸ A review of the literature to date regarding the Ba + O₃ reaction mechanism indicates that it is initiated by long-range electron transfer, that the reaction cross section is ~70A², and that the nascent chemiluminescent products are primarily BaO (A, A'). The reactions of alkaline earth atoms with ClO₂ were studied by Engelke and Zare⁹ in crossed beams chemiluminescence experiments. A particularly fascinating result was the observation of BaCl chemiluminescence from abstraction of the central Cl atom, with formation of O₂ from the outer oxygen atoms of ClO₂ by a concerted mechanism. An additional feature in that system was preference for formation of BaO(A') rather that BaO(A"), even though the exoergicities of both channels were comparable.

In chapter 1, we concluded that the reaction $Ba + NO_2 \rightarrow BaO + NO$ primarily involved long-lived $Ba^+NO_2^-$ collision intermediates even at high collision energies or with electronic excitation of the Ba atom, despite a large reaction exoergicity and relatively few vibrational degrees of freedom in the complex. This behavior was attributed to the thermodynamic stability of the planar $Ba^+NO_2^-$ intermediate and a large exit potential energy barrier for its decay to chemical products. This barrier results from the unfavorable symmetry of the NO_2 2b₁ orbital involved in the electronic rearrangement necessary for decay of $Ba^+NO_2^-$ to doubly ionic $Ba^{2+}O^{2-} + NO_2^{-10}$ Like NO_2^- (E.A.=2.3eV)¹¹, the electron affinities of O_3^- (2.10eV)¹² and ClO_2^- (2.37eV)¹³ are

appreciable, and the possibility of long-range electron transfer will again lead to large reaction cross sections. In the ClO_2 reaction, formation of ground state BaO + ClO products $(\Delta H = -73 \text{ kcal/mole})^{14}$ is of comparable exoergicity to BaO + NO production from the NO_2 reaction $(\Delta H = -61 \text{kcal/mole})^{14}$ (Fig. 1). In the case of ozone, however, formation of $BaO + O_2$ is substantially more exoergic $(\Delta H = -110 \text{kcal/mole})^{14,15}$ In all cases, however, the thermodynamic stabilities of the singly ionic reaction intermediates $Ba^+ClO_2^-$, $Ba^+O_3^-$, and $Ba^+NO_2^-$ are expected to be comparable, 16 lying $\sim 65 \text{kcal/mole}$ lower in energy than the separated neutral reactants (Fig. 2).

Although the chemistry and thermochemistry of ozone¹⁷ and the ozonide anion $(O_3^-)^{18}$ have been studied extensively using a variety of techniques, there have been very few experimental investigations of the dynamics of their chemical reactions under isolated single collision conditions.¹⁹ Analogous studies of ClO_2^{-20} and ClO_2^{-21} have been rare, with substantial uncertainty in the electron affinity only recently resolved.¹³ In this chapter, we present angular and velocity distributions of the products resulting from the reactions of Ba with the 18 electron triatomic ozone and 19 electron chlorine dioxide. A comparison of these reactions with the closely related 17 electron NO_2 will allow us to gain some insight into the general behavior of these molecules.

Although chemiluminescent BaO was the only nascent product observed in previous studies of the gas phase Ba + O₃ reaction, extensive work using matrix isolation techniques with IR or Raman spectroscopy by several groups²²⁻²⁵ has revealed considerably richer chemistry. Andrews and coworkers found that codeposition of Ba and other alkaline earth atoms with O₃²² and O₂²³ in rare gas matrices led to formation

of BaO and BaO₂. In the ozone experiments, BaO₃ was also observed. Similar reactions have also been seen in reactions of alkali atoms²⁵ with O₂ and O₃. As expected, all species were found to be highly ionic owing to the low ionization potentials of the metals and appreciable electron affinities of O,O₂, and O₃. The BaO₂ resulted from several different reaction mechanisms^{22a} due to multiple collisions in the matrix:

$$Ba + O_3 \rightarrow BaO_2 + O \tag{1}$$

Oľ

$$Ba + O_3 \rightarrow BaO + O_2 \tag{2a}$$

$$BaO + O_3 \rightarrow BaO_2 + O_2$$
 (2b)

or

$$Ba + O_2 + M \rightarrow BaO_2 + M \tag{3}$$

Apparently, all three processes were operative, but the relative importance of processes 1 and 2a could not be ascertained. By studying the reactions under single collision conditions in the gas phase, it should be possible to unravel the reaction mechanisms.

The nature of the Ba-O₂ interaction in the gas phase has remained uncertain for many years. ²⁶⁻³⁰ By analogy with the ground state BaO molecule, which is best described in terms of a doubly ionic bond ¹⁰(Ba²⁺O²⁻), we might expect the barium-dioxygen molecule to be considerably more peroxide-like (i.e. Ba²⁺O₂²⁻) than its alkali counterparts which are superoxides (eg. $Cs^+O_2^-$). Although this is certainly true³¹ in the case of solid BaO₂, it has been concluded that BaO₂ formed under matrix isolation conditions is a

superoxide.^{22,23} This conclusion was based on the intraionic O-O vibrational frequency observed for BaO₂ formed in matrix isolation experiments. The observed 1120 cm⁻¹ frequency is substantially higher than that expected for a peroxide (\sim 840cm⁻¹). Based on the matrix isolation work, it was concluded that in the case of the Ba dioxygen molecule, the potential divalency of barium is only seen in the solid. A number of our experimental results on Ba + O₃ provide new insight into the BaO₂ molecule. Based on the recoil velocity of the BaO₂ product, we deduce a lower limit to D_o(Ba-O₂) of 120 kcal/mole. The large binding energy strongly suggests that the molecule does have considerable peroxide character (i.e. Ba²⁺O₂²⁻), in contrast to the much more weakly bound alkali superoxides such as Na⁺O₂⁻ (D_o \approx 40 kcal/mole). Some of the relevant features of the much studied Ba + O₂ reaction, ²⁶⁻³⁰ and the apparent disagreement with the conclusions from the matrix isolation experiments are discussed in this chapter.

Based on the similarities in the molecules NO_2 , O_3 , and ClO_2 (Table 1), we might expect to see many of the same features in the Ba + ClO_2 and O_3 dynamics that were observed in the Ba + NO_2 system. Although at first glance this appears to be true, with direct and long-lived mechanisms contributing to the overall reaction cross section, a considerably larger contribution from a direct mechanism is seen in the O_3 reaction. Whereas the secondary chemically distinct reaction channel $Ba(^1S) + NO_2 \rightarrow BaNO + O$ resulted from a direct mechanism (Chapter 1), in the present reactions both secondary chemical channels, $Ba + ClO_2 \rightarrow BaCl + O_2$ and $Ba + O_3 \rightarrow BaO_2 + O$, result from mechanisms involving long-lived complexes.

Based on the experimental data from the Ba + O₃ experiments alone, a number

of important questions regarding the reaction dynamics could not be answered unambiguously. This complication arose because there was no way to tell for certain which oxygen atoms (apical or terminal) in ozone are responsible for formation of BaO and BaO₂. This was one motivation for the ClO₂ studies since chlorine dioxide possesses many similarities to ozone but the chemically distinct central atom (which leads to chemically distinct products) provides clearer insight into the reaction mechanisms.

II. EXPERIMENTAL

The experimental apparatus employed in this study is identical to that used in the work described in Chapters 1 and 2. The handling of ozone in the context of molecular beams deserves particular mention. Since liquid O_3 is unstable and decomposes to O_2 with possible explosion, condensation of O_3 (bp=-111.9°C)³³ is generally avoided.³⁴⁻³⁶ In our early experiments, the effluent from a commercial ozonizer³⁷ (10% O_3 , 90% O_2) was passed through a pyrex trap containing coarse silica gel cooled to -78°C in a dry ice/acetone mixture. After ~1hour of running time, approximately 10g of deep blue O_3 with a small amount of O_2 was adsorbed on the Si gel. The trap was transported to the molecular beams laboratory and a slow flow of He used to carry the desorbing ozone out of the trap and through the molecular beam nozzle. A thermostatically controlled variable temperature bath was used to slowly warm the O_3 trap in order to maintain a constant concentration as the O_3 was depleted. The O_3 concentration was continuously monitored by its absorption at 280nm³⁸ in a 1cm x 1cm quartz flow cuvette just prior to entry into the molecular beams apparatus. The beam was continuous and contained 10% O_3 , ~1%

O₂, and 89% He at a total pressure of ~300 Torr, after the system was allowed to stabilize for at least 1 hour. With some practice, it was possible to obtain ozone beams of reproducible intensity and velocity over a period of ~6hrs. In our experience, this method is quite safe when several precautions are taken. Pumping undiluted O₃ through rotary pumps is to be avoided, and sealed Si gel traps containing ozone must never be warmed up or pressurized. With daily oil changes, the dilute O₃/He mixture does not seem to seriously harm mechanical pumps. Condensing O₃ in LN₂ foreline traps is not recommended due to the explosive nature of the liquid, especially in the presence of pump oil. It is also important that clean stainless steel, glass, or teflon tubing be used throughout the system since decomposition of O₃ is enhanced by other metals and impurities. Also, ozone rapidly decomposes rubber o-rings. The nozzle tip was heated to $\sim 100^{\circ}$ C to minimize the formation of dimers. Without heating the nozzle, considerable O₅⁺ signal could be observed upon sampling the beam directly with the mass spectrometer. This signal was attributed to fragmentation of $(O_3)_2$ upon electron impact ionization.

In the ozone experiments, the BaO measurements were greatly hampered by the presence of O_2 in the beam since the reaction cross section for Ba + $O_2 \rightarrow$ BaO + O is known to be large. ²⁶⁻³⁰ We found that the quantity of O_2 impurity decreased substantially as the beam was run over a period of several hours, making it nearly impossible to unambiguously distinguish between BaO signal arising from reaction of Ba with O_3 from that with O_2 . With the current apparatus geometry, it is not possible to accurately monitor the fraction of O_2 in the O_3 beam during reactive scattering experiments. A

considerable effort was devoted to finding conditions that would minimize the concentration of O_2 impurity in the O_3 sample. The O_2 impurity could be easily monitored by elastic scattering experiments off Ne. By choosing an appropriate detector angle, it is possible to resolve the separate peaks in the O₂⁺ TOF due to contributions from both O₂ and O₃ elastic scattering. After a considerable amount of experimentation, we concluded that the silica gel itself slowly decomposes O₃, making it impossible to obtain pure O₃ using the silica trap method. We found that the only certain way to obtain a pure O₃/He beam was to employ liquid O₃. Due to the possible explosive nature of O₃₀₀, the sample delivery system and valves were set up to permit remote operation, and the reservoir was clamped behind a safety shield before any O₃ was introduced to the system. As before, approximately 10g of O₃ was deposited on the Si gel trap at -78C. After transporting the trap to the laboratory, the O₃ was condensed into a standard glass bubbler cooled to 77K using He as the carrier gas while slowly warming the trap from - $78 \rightarrow -40$ C over a period of ~ 20 mins. In this way, it was possible to trap 5-10g of the deep blue O₃/O₂ mixture. The bubbler used to trap the liquid O₃ was constructed using a short internal tube so the O₃ level was always below the gas outlet. Great care was exercised to avoid any vibration, shock, or organic impurities, all of which are known to promote explosive decomposition.³⁴⁻³⁶ After the ozone transfer was completed, a slow stream of He was passed into the trap with the effluent directed into the nozzle. The LN₂ trap was carefully replaced with an isopentane slush (-155°C) using remote manipulators, giving an O₃ vapor pressure of ~20Torr.³³ At -155°C, the more volatile³³ O₂ impurity is depleted within several minutes leaving pure O₃. Elastic scattering experiments off He showed a negligible presence of O_2 contamination in the beam when using this method. We found $O_3^+:O_2^+:O^+=4.7:3.0:1.0$, in excellent agreement with the published mass spectral data on pure O_3 sampled from a molecular beam.³⁹ It should be emphasized that this procedure was undertaken with considerable planning and very carefully adopted procedures which could be implemented remotely. In general, the use of liquid ozone is to be avoided due to the hazards associated with its use.

Chlorine dioxide was prepared by the heterogeneous reaction $Cl_{2(g)} + 2NaClO_{2(s)}$ 2NaCl + 2ClO₂ using an adaption of the method of Derby and Hutchinson.⁴⁰ A 10% Cl₂/He mixture was produced by bubbling 1atm He through liquid Cl₂ in a cooled trap. (-78°C) The mixture was directed through a 1" dia. pyrex tube (2' long, wrapped with duct tape) loosely packed with technical grade NaClO₂ (MCB). The effluent, which is nearly pure⁴¹ ClO₂, passed through a U-tube containing P₂O₅ to remove water⁴² and into a trap cooled to -196°C. The Cl₂ impurity was subsequently removed by pumping periodically on the sample while it was warmed slowly to -78°C. The sample was stored in the dark at -196°C. A beam of ClO₂ was formed by bubbling He carrier gas through the trap maintained at -40°C. Our experience is consistent with past reports of spontaneous ClO₂ detonation^{9,43,44} when its vapor pressure exceeds ~80Torr. Extreme care is required when handling ClO₂; toward the end of these experiments explosion of an inline LN₂ cooled trap containing several grams of ClO₂ pointed out the dangers associated with this thermodynamically unstable molecule. No injury or serious damage resulted since extensive precautions had been observed throughout. A recent report¹³ that ClO₂ is "safe" should be taken with caution. An alternative procedure involving in situ production of ClO₂⁴⁵ has been described in the literature. However, in our experience, the method as described always led to the presence of considerable Cl₂ impurity, with the concentration varying from batch to batch. Since the level of this impurity could not be ascertained, an unknown BaCl contribution from Ba + Cl₂ would inevitably result. Thus, the liquid ClO₂ method was found to be preferable—upon starting the beam, the more volatile Cl₂ was depleted rapidly, leaving pure ClO₂. Mass analysis of the beam produced using liquid ClO₂ indicated that it was composed of pure ClO₂ with no indications of impurities such as Cl₂ or Cl₂O. It should be noted that in more recent experiments (Chapter 5), we have found that conversion of Cl₂ to OClO can be nearly quantitative, but only when some moisture is present in the NaClO₂. In that work, the presence of a small quantity of Cl₂ impurity is inconsequential; this was not the case in the present experiment. The nozzle was heated to 150°C to inhibit formation of clusters. We found no evidence for gradual thermal decomposition of ClO₂, even when the nozzle was heated to 150°C, in agreement with the findings of others.⁴¹

III. RESULTS AND ANALYSIS

A. Ba + $ClO_2 \rightarrow BaO + ClO$

The product angular and velocity distributions for the above reaction were recorded at collision energies of 6, 14, and 30 kcal/mole. The Newton diagram for $Ba(^{1}S) + ClO_{2} \rightarrow BaO + ClO$ is shown in Fig. 3 for a nominal collision energy of 14 kcal/mole. The BaO angular distribution (Fig 3.) is forward peaking, indicating the

occurrence of direct reaction mechanisms. However, there is also a wide angle component which we attribute to decay of long lived complexes. Time-of-flight spectra for the BaO product at a collision energy of 14 kcal/mol is shown in Fig. 4. The most notable feature of the data is its similarity to the BaO resulting from the Ba + NO_2 reaction.² We have obtained a reasonably good fit to the data by assuming a single uncoupled $T(\Theta)$ and P(E), (Fig. 5). However, the fit is expected to be improved somewhat through the use of two uncoupled $T(\Theta)$ and P(E) combinations, as employed in the Ba + NO_2 study. This procedure is currently in progress.

B. Ba + $ClO_2 \rightarrow BaCl + O_2$

Nascent BaCl was observed at the parent ion mass and was attributed to the central atom abstraction reaction shown above. The most conclusive evidence for formation of BaCl + O_2 would be observation of the momentum matched O_2 fragment at m/e=32 (O_2^+). Although we did look for the O_2 fragment, the search was unsuccessful. This was expected, based on earlier signal-to-noise calculations. This situation primarily results from high background countrate in the detector at m/e=32 (O_2^+) due to the presence of residual O_2 . A second reason for our inability to detect O_2 results from the kinematic relationship. Due to the heavy BaCl partner, the recoil velocity of the light O_2 is very large. Due to the large center of mass velocity of O_2 (i.e. a large Newton sphere), the signal intensity at the detector is relatively small.

Consequently, we were concerned about the possibility that the signal observed as BaCl⁺ might be attributable to fragmentation of BaClO from the possible BaClO +

O channel, rather than from BaCl + O_2 . This reaction would be analogous to Ba + NO_2 \rightarrow BaNO + O, discussed in Chapter 1. We ruled out this reaction based on the following arguments: 1) BaClO is an unlikely product since it is expected to be thermodynamically unstable⁴⁶ with respect to BaO + Cl; 2) If nascent BaClO had been formed, it is likely to be observable (weakly) at the parent mass or ClO⁺, as was the case for BaNO in Ba + NO_2^2 . Yet no such signal was found; 3) The BaCl product translational energy distribution peaked well away from zero energy as would be expected for a concerted reaction involving central atom abstraction followed by repulsion of BaCl + O_2 ; 4) Nascent BaCl has been observed in past chemiluminescence work.⁹

Zare and coworkers were able to detect nascent BaCl^{+ 9b} (in addition to neutral chemiluminescent BaCl^{+ 9a}), apparently due to reaction 4:

$$Ba + ClO_2 \rightarrow BaCl^+ + O_2 + e^- \qquad (4)$$

This channel is expected to have an even smaller cross section than the chemiluminescent BaCl channel.⁹ We can rule out contribution of nascent ions to our observed signal since our ionizer was floated at +70V. This prevents any positive ions produced in the scattering region from entering the mass spectrometric detector. We also found that the BaCl signal disappeared upon turning off the electron bombardment ionizer, indicating that it does not result from surface ionization of metastable species in the detector.

The angular distribution for the BaCl product from the Ba + ClO₂ reaction at

14kcal/mole is shown in Fig. 6. A good fit of this data and the BaCl TOF (shown in Fig. 7), was obtained using a single uncoupled P(E) and T(Θ) combination. The best fit to this data at 14kcal/mole is shown in Fig. 8. The angular distribution was found to exhibit forward-backward symmetry in the center-of-mass frame of reference at all collision energies.

The yield for the BaCl + O_2 channel was found to be approximately 10% that of the BaO + ClO channel at all collision energies studied. This quantity was calculated from the optimized P(E) and T(θ), after normalization of the observed signal intensities. In the calculation, we assumed equal detection efficiencies for BaO and BaCl at their parent ion masses. This is a reasonable assumption in the present case since it is known that the parent ion peaks for both BaO and BaCl are comparable to the daughter ion peak at Ba⁺. The fragmentation patterns are also known to be relatively insensitive to internal excitation of the parent neutral molecule.² The ionization cross sections for both species are calculated to be nearly identical.⁴⁷

C. Ba + $O_3 \rightarrow BaO + O_2$

The product angular and velocity distributions were recorded at three collision energies between 6 and 40 kcal/mole using liquid ozone. The BaO angular distribution obtained at a collision energy of 14 kcal/mole was more forward peaking than in the case of Ba + NO₂ or Ba + ClO₂. Although the BaO angular distributions clearly indicate a substantially larger yield of forward scattered products than in the Ba + NO₂ and ClO₂ systems, a wide angle component is seen. The time of flight data at this collision energy

is shown in Fig. 9; the translational energy and CM angular distribution resulting from the data is presented in Fig. 10. A BaO product flux contour diagram, superimposed on the nominal Newton diagram is depicted in Fig. 13. By analogy with the NO₂ system, the involvement of long-lived Ba⁺O₃ collision complexes is a logical explanation for backscattered BaO. However, an alternative mechanism that was impossible for Ba + NO₂, involving abstraction of the central O atom from O₃ resulting in O₂ by formation of a chemical bond between the outer oxygen atoms in ozone, is possible in principle. It is likely to be dominated by small impact parameter collisions with near C_{2v} approach of the Ba atom towards the apical oxygen atom in ozone, possibly leading to backscattering of the BaO molecule. An analogous process has been postulated by van den Ende and coworkers 19a to account for a sharply backscattered NO2 peak in the reaction NO + $O_3 \rightarrow NO_2 + O_2$. If such a reaction were operative in the present reaction, analysis of our scattering data using an iterative forward convolution approach might best involve two separable $T(\theta)$ and P(E) combinations—one involving a forward scattered "stripping" channel and the other due to abstraction of the central O atom with BaO backscattering. However, we have chosen not to use this approach. Although the data presented here was analyzed using a single P(E) and $T(\theta)$ combination, further analysis in progress employs the same model used in the analysis of the BaO from the Ba + NO₂ data, with one forward-backward symmetric channel and a second forward scattered angular distribution. This approach is largely inspired by our observations in the analogous system Ba + ClO₂ where a substantial yield of BaCl + O₂ was observed, but the best fit BaCl angular distribution was forward-backward symmetric rather than

backscattered. We believe that central atom abstraction from ozone, if it were to occur at all, will likely show a forward-backward symmetric BaO angular distribution. (This will be discussed in detail in section IV-A.) The fact that the BaO₂ product angular distribution exhibited forward-backward symmetry clearly indicates the existence of a stable Ba $^+$ O₃ intermediate which will likely branch into both BaO + O₂ and BaO₂ + O. We cannot envision any physically realistic mechanism requiring us to incorporate a separate reaction channel involving BaO backscattering.

D. Ba + $O_3 \rightarrow BaO_2 + O$

The formation of nascent BaO_2 from a single encounter between $Ba + O_3$ was clearly observed. Product velocity and angular distributions were recorded at three collision energies. Since it is not possible that O_2 contamination can lead to observation of BaO_2 at the detector, as was verified in experiments using pure O_2^{48} , extensive experiments using liquid ozone were deemed to be unnecessary and the data presented here employed O_3 directly from the silica gel trap. However, an experiment was carried out using liquid O_3 to verify that the BaO_2 signal resulted exclusively from reaction with O_3 , and was not affected by the presence of O_2 impurity in the O_3 beam.

There is no way to precisely determine the BaO₂ fragmentation patterns upon electron bombardment ionization since the signal observed at BaO⁺ is strongly dominated by nascent BaO from the BaO + O₂ channel. Analysis of the Ba⁺ data is complicated by the fact that it results from fragmentation of both BaO₂ and BaO upon electron bombardment ionization, as well as from nonreactive scattering events (although this is

probably minor). The data presented here were recorded with the mass spectrometer tuned to $BaO_2^+(m/e=170)$. We believe that the internal energy dependence of the BaO_2 fragmentation pattern is likely to be small. This is based on the large BaO_2 binding energy (this work), as well as a large binding energy for BaO_2^+ . Good fits to the BaO_2 product TOF (Fig 11) and angular distributions were obtained using a single uncoupled P(E) and $T(\theta)$ combination (Fig 12). The center of mass angular distribution exhibited forward-backward symmetry at all collision energies. The BaO_2 product flux contour map, superimposed on a nominal Newton diagram at 14 kcal/mol is shown in Fig. 13.

IV. DISCUSSION

A. Ba + ClO,

There is remarkable similarity between the BaO resulting from reaction of barium with ClO₂ with that from NO₂. Again, one mechanism involves long-lived complexes and a second involves direct reactions. Perhaps the only substantial difference is that the average translational release for the present reaction is somewhat less, despite a slightly larger reaction exoergicity. Although we have not attempted to measure relative reaction cross sections for Ba + NO₂ and ClO₂, they are expected to be comparable since the electron affinities of the molecules are nearly identical. The BaO + ClO channel is dominant and accounts for ~90% of the total reaction cross section at all collision energies.

As was the case for NO₂, formation of the negative ion fragments from ClO₂ + e is generally endoergic with the notable exceptions of the three-center rearrangements

forming $Cl^- + O_2$ or $Cl + O_2^-$:50

ClO₂ + e⁻
$$\rightarrow$$
 ClO₂⁻ \rightarrow ClO⁻ + O = +11.5 kcal/mole (5)
ClO₂ + e⁻ \rightarrow ClO⁻ + O = +11.5 kcal/mole (6)
 \rightarrow ClO + O⁻ = +26.2 kcal/mole (7)
 \rightarrow Cl⁻ + 2O = +38.7 kcal/mole (8)
 \rightarrow Cl + O₂⁻ = -5 kcal/mole (9)
 \rightarrow Cl⁻ + O₂ = -79.5 kcal/mole (10)

There appears to be no information regarding cross sections for these processes. However, electron attachment to ClO₂ at low energy should lead to formation of a long-lived anion, ⁵⁰ due to a large potential energy barrier for the concerted molecular rearrangements (processes 9,10). ^{53b} The energetics of ClO₂ decomposition are shown in Fig. 14. Based on the thermodynamics, the situation is completely analogous to NO₂ and its anion: the neutrals have large electron affinities, and the anion is thermodynamically stable with respect to fragmentation. ^{2,50,51} For most collision trajectories, long-range electron transfer results in formation of a long-lived collision intermediate. The highest unoccupied molecular orbital of ClO₂ is ⁵² (2b₁)¹. Long-range electron transfer into the half-occupied 2b₁ orbital leads to ClO₂ (¹A₁). ^{9,49} Whereas electron attachment to NO₂ results in considerable decrease in bond angle, (from 133.9° to 117.5°), a much smaller change from 117° to 111° (Ref. 58b, Table 1) occurs in the case of ClO₂. However, the O-O bonding character of the 2b₁ orbital and the large exoergicities for the BaO₂ + Cl and BaCl + O₂ channels suggest that concerted

decomposition might occur. Although little information is available regarding the next unoccupied orbitals of ClO₂, there is good evidence that the next higher orbital is of Rydberg character.⁵² Like the Ba + NO₂ reaction, electron transfer leads to symmetric stretching and bending in the ClO₂ ion. Since the reaction coordinate for formation of BaO + ClO involves asymmetric stretching, a substantial fraction of the collisions result in formation of long-lived complexes. The potential energy barrier associated with orbital rearrangement required for formation of ground state doubly ionic BaO + ClO products, and the barrier for the concerted process favor formation of long-lived complexes.

The BaCl + O_2 reaction is of particular interest since the reaction has been found to involve concerted central atom abstraction. Although such reactions are extremely rare under single collision conditions, we find that the yield of BaCl + O_2 is appreciable (~10%). Engelke and Zare concluded that long-range electron transfer into the $2b_1$ orbital led to a decrease in ClO_2 bond angle with enhanced bonding between the oxygen atoms resulting in formation of BaCl + O_2 . Based on the observation that the internal energy distribution was nonstatistical, they concluded that formation of chemiluminescent BaCl resulted from a direct reaction mechanism, rather than from one involving long-lived intermediates. Although the large BaCl signal level observed in our experiment is consistent with a mechanism initiated by long-range electron transfer into the $2b_1$ orbital, the nearly perfect forward-backward symmetry observed in the product angular distribution is very strong evidence that the $Ba^+ClO_2^-$ intermediate undergoes at least several rotations before formation of most BaCl product molecules. The existence of an

osculating complex with lifetime of about one rotational period is ruled out based on the insensitivity of the angular distribution to collision energy. The nonstatistical internal energy distribution observed by Zare and coworkers thus might be indicative of different reaction mechanisms for formation of chemiluminescent and ground state BaCl. Alternatively, the internal energy distribution that they observed might result from the effects of a large potential energy barrier in the BaCl $+ O_2$ exit channel.

We find that the yield of BaCl relative to BaO is insensitive to collision energy. One possible reaction mechanism would involve a common Ba⁺ClO₂⁻ intermediate with kinetic competition for its decay to BaO + ClO or BaCl + O₂. The lack of a collision energy dependence could be taken as evidence that the potential energy barriers for the two processes are comparable. Alternatively, the observations might be interpreted in terms of two different reaction mechanisms arising from different reactant approach geometries. Since both apparently involve long-lived intermediates, this would suggest the participation of two different geometric isomers of Ba⁺ClO₂⁻ with formation of BaCl + O₂ resulting from a Ba-Cl bonded complex and BaO + ClO from decay of a bidentate structure analogous to that postulated in the Ba + NO₂ reaction (Chapter 1). Matrix isolation experiments have indicated that the Cs⁺ClO₂⁻ complex has C_{2v} symmetry.^{53,54} Although the authors mentioned a bidentate structure, it appears that this assignment was based only on the apparent analogy with similar molecules such as Cs⁺NO₂, Cs⁺CO₂, or Cs⁺SO₂, all of which are believed to be bidentate. However, in the case of Cs⁺ClO₂ or Ba⁺ClO₂, a monodentate C_{2v} structure should be substantially more favorable than in the case of NO₂ or CO₂. This is due to the stronger interaction between the metal cation and the central atom in ClO₂ than in NO₂ or SO₂, because of the greater electronegativity of the Cl atom.

It is interesting to note that the most exoergic reaction channel, Ba + $ClO_2 \rightarrow$ $BaO_2 + Cl$ is not observed. Based on Fig. 14, formation of $Cl + O_2$ from decay of ClO₂ is nearly thermoneutral, whereas formation of Cl⁻ + O₂ is highly exoergic. Of course, this is due to the much greater electron affinity of Cl. Although both concerted channels will have substantial potential energy barriers, based on the simple energetics of the anions we would expect that formation of BaCl will be more energetically favorable than BaO₂. It should also be noted that the ground state of BaO₂ correlates to a doubly ionic (excited state) of BaClO₂, whereas the BaCl molecule is only singly ionic. Consequently, formation of BaCl from concerted decomposition of a monodentate complex is expected to involve substantially less electronic rearrangement than a hypothetical BaO₂ product from a bidentate intermediate. Also, formation of BaCl or BaO₂ necessitates considerable deformation of the BaClO₂, likely requiring large amplitude symmetric bending of ClO₂. As mentioned previously, electronic excitation into the 2b₁ orbital will lead to "scissoring", i.e. excitation of the reaction coordinate. Although this is very favorable in the Ba-Cl bonded intermediate, in a bidentate BaClO₂ molecule, it is likely that such motion will be "damped" considerably by the presence of In light of these considerations, we are not surprised by the absence of the BaO_2 channel in the $Ba + ClO_2$ reaction.

There has been extensive discussion in the literature^{44,55-59} regarding photochemical

isomerization of the symmetric ClO₂ molecule (i.e. OClO) to Cl-OO. Electronic excitation of ClO₂ corresponds to excitation of an electron into the 2b₁ orbital; the same orbital populated by long range electron transfer from Ba in the Ba + ClO₂ reaction. Thus, it might be argued that the gas phase Ba + ClO₂ reaction could involve a mechanism involving isomerization of the ClO₂ in BaClO₂ to the peroxy form, followed by simple bond rupture resulting in BaCl + O₂. However, this mechanism would not be consistent with the measured BaCl translational energy distribution. The P(E) clearly peaks well away from zero, as expected for a reaction having a substantial potential energy barrier in the exit channel. An isomerization mechanism followed by simple bond rupture forming BaCl should not have an appreciable exit potential energy barrier, and should result in a P(E) peaking near zero energy. Indeed, as will be pointed out in Chapter 5, the hypothesized gas phase isomerization mechanism is in fact not correct for an isolated electronically excited ClO₂ molecule.

B. Ba + $O_3 \rightarrow BaO + O_2$

As was the case for simple bond rupture of nitrite (NO₂) and chlorite (ClO₂), dissociation of the ozonide ion (O₃) to O⁻ + O₂ is more endoergic than is formation of O + O₂ from the neutral molecule. However, it is important to note an important feature of the energetics that is unique to O₃. We postulate that this one feature of ozonide has a substantial effect on the reaction dynamics. Whereas O⁻ production from dissociative electron attachment to NO₂ and ClO₂ is endoergic, the process O₃ + e \rightarrow O₂ + O⁻ is appreciably exoergic^{14,60,61} (Fig. 14). This is reminiscent of the behavior seen

in the halogens.⁶² Indeed, in low energy attachment experiments, both O₃⁶⁰ and Br₂⁶² possess finite cross sections for dissociation even at zero electron kinetic energy. This suggests that the Ba + O₃ reaction might show features which are more analogous to the alkali + halogen system which has been interpreted using the spectator stripping mechanism.^{63,64} In spectator stripping, long-range electron attachment leads to immediate dissociation of the anion with forward scattering of the metal containing molecule. This hypothesis appears to be supported by the angular and velocity distributions that we have obtained for the BaO product.

In the spectator stripping limit, the atom (X) not involved in formation of the chemical bond in the reaction $M + X_2 \rightarrow M^+X^- + X$ is assumed to remain unperturbed throughout the encounter. The magnitude and direction of its final velocity vector is identical to that of the incident X_2 molecule. It might appear inappropriate to extend such a simple mechanism to a reaction of a triatomic molecule, since the molecular fragment (O_2) possesses vibrational and rotational degrees of freedom. However, this limiting case is likely to be approached for a substantial fraction of all reaction trajectories in the present reaction for several reasons: The ozone molecule and ozonide anion are extremely weakly bound; dissociation of the anion is likely to be facile in the field of Ba^+ . Even more importantly, the resulting molecular fragment (O_2) is strongly bound, but has a bondlength only 0.07\AA^{65} shorter than that of the neutral ozone molecule.

The calculated BaO-O₂ recoil velocity based on the spectator stripping model leads to a mean product translational energy of 10kcal/mole, and the spectator stripping

laboratory angle for the BaO product is 15 degrees. The small release of translational energy with large internal excitation of the product molecules, and forward peaking angular distribution, is consistent with a substantial contribution from a spectator stripping mechanism.

Formation of electronically excited BaO will most likely result from long-range electron transfer followed by immediate formation of Ba⁺O⁻ + O₂. The singly ionic structure of BaO is electronically excited.¹⁰ Although the angular distribution of the chemiluminescence has not been measured precisely, it was relatively easy for us to observe this emission by eye after turning off the lights and becoming accustomed to the dark. A long (~30cm) chemiluminescent streak could be observed inside our apparatus; apparently this resulted from emission by the relatively long-lived BaO (A') state.¹⁰ A particularly interesting feature of the chemiluminescence is that it appeared to be very nearly confined to the Ba beam axis. This indicates that it came from a direct mechanism with forward scattering of the electronically excited BaO molecule with low recoil energy. This appears to be consistent with the spectator stripping mechanism leading to electronically excited products.

Despite the fact that formation of ground state products is ~50 kcal/mole more exoergic than the Ba + NO₂ reaction, we observe smaller release of translational energy in the present reaction, indicating that the products are highly internally excited. Since the single collision yield of electronically excited BaO is known to not exceed 20%, it is certain that a large fraction of products are $O_2(^1\Delta)$ or $O_2(^1\Sigma)$. Indeed, formation of ground state BaO($^1\Sigma$) from the singlet reactants requires formation of electronically

excited singlet O_2 to conserve spin. The requirement of spin conservation in the context of barium atom reactions has received considerable attention in the past. Zare⁶⁶ and Herm⁶⁷ have both studied the reaction

$$Ba(^{1}S) + SO_{2}(^{1}A_{1}) \rightarrow BaO(^{1}\Sigma^{+}) + SO(^{3}\Sigma^{-}). \tag{11}$$

At low collision energies, the only thermodynamically possible reaction is spin forbidden since the spin allowed process:

$$Ba(^{1}S) + SO_{2}(^{1}A_{1}) \rightarrow BaO(^{1}\Sigma^{+}) + SO(^{1}\Delta)$$
 (12)

is endoergic by 14.1 kcal/mole. However, strong BaO signal was observed and resulted from decay of long-lived Ba⁺SO₂⁻ collision complexes. The existence of collision complexes in that reaction was not surprising in view of the small reaction exoergicity, and earlier indications⁶⁸ that long-lived Cs⁺SO₂⁻ complexes were involved in nonreactive scattering of Cs + SO₂. The very large cross section for the spin forbidden process (>50Å) was attributed to a mechanism involving intersystem crossing of the initially formed singlet intermediate:^{66,67}

$$Ba(^{1}S) + SO_{2}(^{1}A_{1}) \rightarrow {^{1}}(Ba^{+}SO_{2}^{-}) \rightarrow {^{3}}(Ba^{+}SO_{2}^{-}) \rightarrow BaO(X^{1}\Sigma^{+}) + SO(^{3}\Sigma^{-})$$
 (13)

Enhanced spin-orbit coupling due to the presence of the heavy Ba and S atoms apparently

leads to an appreciable rate of intersystem crossing in this system. Since the lifetime of the intermediate complex is greater than several picoseconds, with the only alternative process being decay back to reactants, the probability for the spin forbidden process in collision complexes can be large. A similar behavior in the analogous ozone reaction, with formation of spin forbidden ground state $BaO(^{1}\Sigma^{+}) + O_{2}(^{3}\Sigma^{-})$, also appears feasible due to the effect of the heavy Ba atom. However, in the present case, our product angular distributions indicate that the dominant reaction mechanism involves a direct process, with forward scattering dominating at all collision energies. This implies that most products are formed on a much shorter average timescale in this reaction. In reactions of light atoms, a short lifetime would suppress spin forbidden processes. For an atom such as Ba, however, this is probably not a major consideration. It is worth noting, however, that based on the low average translational energy we observe in the BaO products, a large fraction of the product O₂ molecules must be electronically excited. Since the dominant BaO is $X^1\Sigma^+$, a large fraction of the BaO + O_2 products do result from reactions which conserve spin.

We have already mentioned that the backscattered BaO molecules could result from direct abstraction of the central oxygen atom with formation of O_2 from the outer oxygen atoms in ozone. Unfortunately, we see no unique features in our data that can be associated with this process. Based on our observations for Ba + $ClO_2 \rightarrow BaCl + O_2$, even if such an abstraction were to occur in the Ba + O_3 reaction, it would likely lead to a forward-backward symmetric angular distribution. Although we do find that the BaCl product from the concerted channel peaks well away from zero energy, the maximum

energy remains very low. Clearly, most of the energy is channeled into internal excitation of the products. If central atom abstraction were to occur in the Ba + O₃ reaction, the product is likely to be hidden under the BaO from reaction of the outer oxygen atoms resulting from decay of Ba $^+$ O₃ intermediates.

C. Ba + O_3 -> Ba O_2 + O

One of the most intriguing observations in the rather extensive history of Ba oxidation studies $^{26\cdot30}$ was that BaO (A $^{1}\Sigma^{+} \rightarrow X^{1}\Sigma^{+}$) emission above 17588cm $^{-1}$ can be seen from the Ba + O $_{2}$ reaction despite a reaction exoergicity for Ba + O $_{2} \rightarrow$ BaO + O of only 4547cm $^{-1}$. Beginning with the earliest experiments, there was considerable discussion and speculation regarding a likely multi- step mechanism consistent with sufficient energy pooling to permit a high pressure chemiluminescence yield approaching 10%. Although the common denominator in all mechanisms has been the participation of BaO $_{2}$, this species has proven to be particularly elusive to direct observation. The clearest evidence for its transient existence was obtained by Herm and coworkers in the forward-backward symmetric BaO angular distribution observed in crossed beams studies of the Ba + O $_{2}$ reaction. Their work indicated that the lifetime of the BaO $_{2}$ intermediate exceeded several rotational periods (several psecs).

Edelstein and coworkers^{30a} found a series of bands in diffusion flame experiments that they attributed to emission from electronically excited BaO₂ produced in three-body recombination reactions. These so called "Y-bands" appeared near 700nm. Although more recent experiments by Reuther and Palmer^{29b} provided additional support for their

hypothesis, the actual nature of the transitions leading to the emission remains unknown, and no thermodynamic or structural information exists on gas phase BaO_2 to date. Although the existence of long-lived BaO_2 collision complexes even at collision energies of 40kcal/mole suggests that $D_o(Ba-O_2)$ is likely to be substantial, Herschbach^{27c} has suggested that a long intermediate lifetime might be attributable to hopping between singlet and triplet surfaces rather than from thermodynamic stability alone. Other comments that $D_o(Ba-O_2)$ is likely to be substantial are scattered throughout the literature, ²⁶⁻³⁰ but a definitive measurement does not exist.

The Ba-O₂ bond dissociation energy was obtained from the maximum translational energy release in the reaction Ba + O₃ \rightarrow BaO₂ + O. Using the same arguments presented in the determination of D_o(Ba-NO) in the chapter 1, the following relation may be derived:

$$D_o(Ba-O_2) \ge E_{trans \ max, BaO2/O} + D_o(O-O_2) - E_{coll} - E_{int,O3}$$
 (14)

Based on the results of the lowest collision energy experiment: $E_{trans\ max,\ BsO2/O}=110$ kcal/mole, $E_{coll}=16$ kcal/mole and $E_{int,\ O3}=1$ kcal/mole we obtain $D_o(Ba-O_2)\geq 120$ kcal/mole. Since both reactants are singlets, spin conservation requires formation of triplet BaO₂ if the O atom is in the ground state (³P). Although we will argue that the ground state of BaO₂ is a singlet, by analogy with the Ba + SO₂ reaction, the involvement of long-lived complexes in formation of BaO₂ is expected to lead to relaxation of the requirement of spin conservation. Thus at least some BaO₂ + O

products will be formed in their ground electronic states and equation 14 will an equality to within the expected error limits of \pm 20 kcal/mole.

Although there are no experimental or theoretical values available for alkaline earth-dioxygen bond strengths, considerable insight into the nature of the interaction can be obtained by comparison with the analogous quantity for the alkali superoxides which are at most singly ionic due to those metals' monovalent nature:

Molecule	\underline{D}_{o} (M-O ₂) kcal/mole ⁷⁰⁻⁷²
Li-O ₂	52 ± 5
Na-O ₂	39 ± 5
K-O ₂	41.5 ± 5
Rb-O ₂	43 ± 5
Cs-O ₂	44.5 ± 5

These values are in very good agreement with bond strengths predicted based on a simple Hess cycle⁷² for singly ionic molecules:

$$D_o(M-O_2) = D_o(MO-O) + D_o(MO) - D_o(O_2)$$
 (15)

Since the oxygen-oxygen vibrational frequency in the alkali superoxides is nearly identical to that of the free superoxide ion, it is customary⁷² in these calculations to take $D_o(MO-O) \approx D_o(O-O^-) = 4.09eV$. Since $D_o(O_2) = 5.114 eV$, the dissociation of the

singly ionic metal superoxide is related to that of the monoxide by:

$$D_o(M-O_2) = D_o(MO) - 1.02 \text{ eV}.$$
 (16)

If BaO₂ is a superoxide (singly ionic), we expect that its bond dissociation energy would be well approximated by the above relationships. Failure to observe good agreement with a value based on a singly ionic structure could be taken as evidence for substantial doubly ionic character. The ground state BaO($X^1\Sigma^+$) molecule ($D_o = 5.77eV$) is best described as Ba²⁺O²⁻ (Ref 10), whereas the first excited state A($^1\Pi$) at 2.17eV^{73,74} is well represented as Ba⁺O^{-10,74} Indeed, Langhoff and Bauschlicher⁷⁴ have pointed out that $D_o(MO^*) - D_o(MF)$ for the excited state A($^1\Pi$) alkaline earth oxides and the ground state alkaline earth fluorides is nearly constant at 2.4eV. This is good support of a singly ionic interaction for both species. Thus, for singly ionic BaO, $D_o(Ba-O) = 5.77-2.17 = 3.6 \text{ eV}$. Assuming a superoxide structure, we should expect that $D_o(Ba-O_2)$ should lie 1.02 less than this, at about 2.6eV (60 kcal/mole).

The experimental Ba-O₂ binding energy obtained from this experiment is twice that predicted assuming a superoxide structure. This is good evidence that the BaO₂ resulting from the Ba + O₃ reaction does have substantial peroxide-like character. It is interesting to note that D₀(Ba-NO) = 65kcal/mole (Chapter 1) is very close to the value calculated for singly ionic BaO₂. This is not surprising since all available evidence indicates that alkaline earth nitroxides are well represented as M⁺NO (Chapter 1). By analogy with the alkaline earth monoxides, we expect electronic excitation of ground

state barium peroxide to be largely charge-transfer in nature, with the excited state well represented by Ba^+O_2 . The emission attributed to BaO_2 by Edelstein^{30a} and coworkers at 650nm suggests the existence of two BaO_2 electronic states separated by approximately 1.9eV. This is somewhat less than the 2.6 eV (60kcal/mole) gap we have calculated between our experimental bond energy for the peroxide and that estimated for the superoxide. However, this difference is perhaps not too surprising based on the crude approximations involved here. If we take $Ba + O_3$ to be at zero energy, it appears very reasonable that the ground state of BaO_2 , represented as $Ba^{2+}O_2^{2-}$ lies at approximately - 120kcal/mole, and the excited charge transfer state ($Ba^+O_2^-$) at an energy 1.9 eV higher than this, near -75kcal/mole.

Our finding that the ground state of gaseous BaO₂ has substantial peroxide character is consistent with the known electronic structure of the solid³¹ but is somewhat contradictory to the results of matrix isolation experiments. The matrix conclusions^{23a} were based on the observation of a BaO₂ intraionic O-O stretching frequency of 1120cm⁻¹, as expected for a superoxide molecule. It appears that the discrepancy may be reconciled in light of our work on the reaction dynamics of Ba with oxygen containing species. The singly ionic (electronically excited) species is likely to be metastable in the matrix due to the presence of a large potential energy barrier between singly and doubly ionic states. However, collisional processes should induce internal relaxation of at least some of the BaO₂ molecules to the ground electronic state. Thus, at least some ground state BaO₂ should have been observable. Unfortunately, no mention is made in the original papers as to the absence or presence of BaO₂ absorptions in the peroxide

stretching region. Presumably this is because a BaO₂ molecule possessing a 1120cm⁻¹ frequency had been identified, and appeared quite analogous to earlier observations of other molecules such as Cs⁺O₂. Apparently, the possibility for other BaO₂ frequencies was not considered. However, one IR absorption in the peroxide stretching region at 752cm⁻¹ remained unassigned and in principal might be consistent with Ba²⁺O₂²⁻. The difference between this value and that in solid BaO2 is perhaps not unreasonable due to the effects of their different environments. The 752 cm⁻¹ absorption was postulated to be due to a hypothetical Ba₂²⁺O₂² species with a C_{2v} nonplanar structure. However, in view of the known strength of the BaO bond, electronic rearrangement with formation of two BaO molecules (even electronically excited) or a (BaO)₂ dimer structure (having a BaO-BaO binding energy of ~80kcal/mole^{22,23}) should be highly exoergic. Unless some extremely large potential barrier existed for decomposition of Ba₂²⁺O₂², its existence seems unlikely. It is also interesting to note that concentration dependence studies described in the original work actually supported a 1:1 Ba:O₂ stoichiometry for the unassigned line, again consistent with Ba2+O22- rather than Ba22+O22-.

An alternative explanation for the failure to observe Ba²⁺O₂²⁻ in the matrix is that its infrared absorption is very weak. This is quite likely in view of the small change in dipole moment expected for O-O stretching in a triangular BaO₂ molecule. Raman spectroscopy was used in the study³¹ of solid BaO₂, and might also be a sensitive probe in the matrix.

Gas phase and matrix environments can lead to vastly different chemistry in the same systems. Indeed, one of the powers of the matrix isolation technique is the ability

to observe weakly bound molecules, which in the matrix remain trapped in local potential energy minima. In the gas phase, most thermodynamically unstable species undergo unimolecular decomposition or reaction within several rotational periods or less. The gas phase $Ba + O_2$ reaction is initiated by formation of the singly ionic $Ba^+O_2^-$ intermediate. Under collision free conditions, it must decay within several picoseconds, either back to reactants or to the only thermodynamically accessible products, BaO ($^1\Sigma$) + $O(^3P)$. Radiative processes are far to slow to stabilize the intermediate. However, in the matrix, some of the $Ba^+O_2^-$ species can become stabilized as their vibrational energy is transferred to the host matrix molecules which have an effective temperature of only a few Kelvin. Subsequently, there will be insufficient energy available for decay back to $Ba + O_2$ or to the more thermodynamically stable $Ba^2^+O_2^{-2}$. In the matrix reaction, the product of the initial "harpooning step" may become trapped, and can be observed in the infrared spectrum.

Many of the other observations made over the last 20 years related to BaO_2 can be rationalized in terms of a substantial potential energy barrier between superoxide and peroxide electronic structures. However, in the gas phase $Ba + O_2$ reaction, the situation is even more complex. The reaction is known to result from harpooning, with formation of a long-lived complex. However, because the $Ba(^1S) + O_2(^3\Sigma)$ reactants approach on a triplet surface; the initially formed complex is expected to be a singly ionic triplet molecule. This reaction is known to involve a complex even at collision energies approaching 50kcal/mole. Some authors have taken this to indicate the existence of a very strong BaO_2 intermediate, which is confirmed in the present work. However,

the strongly bound intermediate is a peroxide. Based on the molecular orbitals of O₂, the strongly bound barium peroxide molecule will be a singlet. Thus, the long lifetime in the Ba + O₂ reaction does not derive from the thermodynamic stability of the initially formed Ba⁺O₂ intermediate, since its bond energy does not greatly exceed 75 kcal/mole. The potential energy barriers that lie along the reaction path will play an important role. The reaction path involves hopping from the singly ionic triplet surface to the strongly bound doubly ionic surface. This facilitates the electronic rearrangement necessary for formation of the doubly ionic ground state $BaO(^{1}\Sigma)$, the only thermodynamically accessible product. However, thermodynamics also dictates formation of O(³P). Thus, a final hop back to the triplet surface will be required for exit. Although the ground state of BaO₂ does represent a very deep minima, it is not accessed in the initial harpooning step but is ultimately accessed in the reaction, since no other channels (other than decay back to reactants) exist. We conclude that the long reaction timescale in the Ba + O₂ reaction results from both factors-- multiple surface hopping and a strong BaO₂ intermediate.

V. CONCLUSIONS:

We have found that although the gas phase chemistry of Ba + O₃ and Ba + ClO₂ is highly complex, most of our observations can be rationalized in terms of rather fundamental principles. These principles derive from the chemistry and thermochemistry of the anions, and from the same molecular orbital concepts which have proven so useful in understanding general trends in triatomic molecules. As we saw in the NO₂ reaction,

the initial features in reactions involving divalent Ba are similar to those of monovalent species. However, the participation of the second valence electron does lead to additional features in the reaction dynamics when doubly ionic products are formed. Since this divalent character, which leads to the large reaction exoergicities is not seen in the initially formed singly ionic intermediate, the reaction dynamics are often controlled by the dynamics of the unimolecular decomposition of a long lived complex. The participation of reaction complexes in general leads to product angular distributions exhibiting forward-backward symmetry, which unfortunately reduces the potential insight into favorable reaction geometries, often afforded in direct reactions. However, the present technique is uniquely powerful in many respects. Not only can we identify all chemical products without the handicap of having to know their spectra, but we can gain insight into the dominant features of the potential energy surface which will control the breakup of singly ionic species into their more stable divalent products.

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Table I: Molecular Parameters for $NO_2,\ O_3,\ ClO_2$ and Their Anions.

Neutral	R(X-O) (Å)	$ heta_{ m oxo}$ Ani (deg)	ion	R(X-O) (Å)	$\theta_{\text{oxo-}}$ (deg)
NO ₂	1.1938ª	133.857ª	NO ₂ ·	1.252*	117.5ª
O_3	1.2717°	116.78°	O ₃ -	1.3415 ^b	112.6 ±2 ^b
ClO ₂	1.47°	117°	ClO ₂ -	1.70°	110°

^{*}Ref 11 *Ref 12b *Ref 12c

dRef 44

Ref 50b

VII. FIGURE CAPTIONS

- 1. Energetics of the Ba + ClO₂ reactions. Energy levels for the thermodynamically allowed reaction channels are shown. The Ba⁺ClO₂ is estimated to lie at approximately -65kcal/mole with respect to Ba + ClO₂. See text for details.
- 2. Energetics of the Ba + O_3 reactions. Energy levels for the thermodynamically allowed reaction channels are shown. The Ba⁺O₃⁻ is estimated to lie at approximately -65kcal/mole with respect to Ba + O₃. See text for details.
- 3. BaO laboratory angular distribution for Ba + ClO₂ → BaO + ClO at nominal collision energy of 14 kcal/mole. Angular distribution is very similar to that seen from Ba + NO₂. Effect of laser excitation on reaction dynamics will be discussed in full paper.
- 4. BaO TOF spectra from Ba + ClO_2 reaction at $E_{coll} = 14$ kcal/mole.
- 8. Relative product translational energy distribution P(E) and center of mass angular distribution $T(\theta)$ for the BaO channel from Ba + ClO₂ reaction. A single uncoupled $T(\theta)$ and P(E) was used in the analysis.
- 6. BaCl laboratory angular distribution from Ba + ClO_2 at $E_{coll} = 14$ kcal/mole. Effect of laser excitation on reaction dynamics will be discussed in full paper.
- 7. BaCl TOF spectra from Ba + ClO_2 reaction at $E_{coll} = 14$ kcal/mole. A single uncoupled P(E) and T(θ) combination was employed.
- 8. P(E) and $T(\theta)$ for the BaCl channel from Ba + ClO₂ reaction.
- 9. BaO TOF spectra from Ba + O_3 reaction at $E_{coll} = 14$ kcal/mole.

- 10. P(E) and $T(\theta)$ for the BaO channel from Ba + O₃ reaction.
- 11. BaO₂ TOF spectra from Ba + O₃ reaction at 14 kcal/mole.
- 12. BaO₂ Product translational energy distribution and center of mass angular distribution from reaction at 14 kcal/mole
- 13. BaO_2 and BaO product flux contour maps superimposed on nominal Newton diagrams for $Ba + O_3$ reaction.
- 14. Energetics of ClO₂, O₃ and its decay products. See text for details.

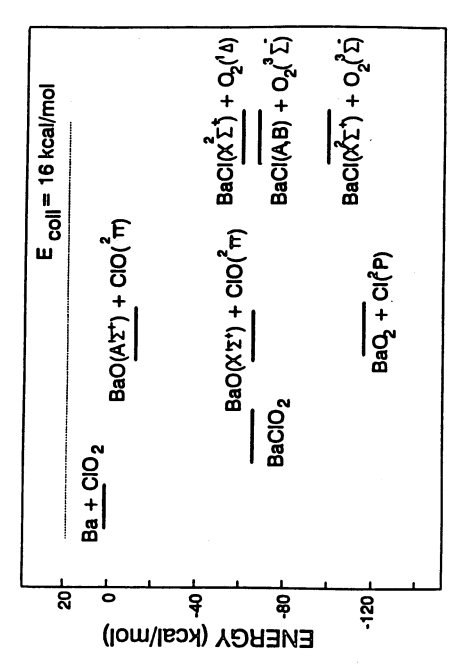


Fig. 1

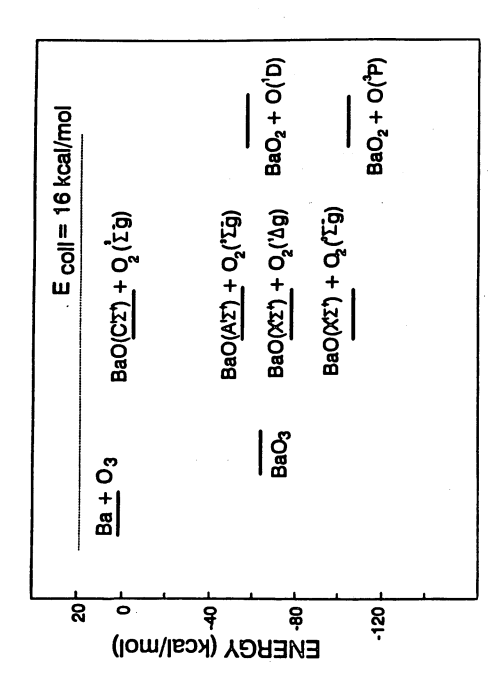


Fig. 2

Ba + ClO2 - BaO + ClO

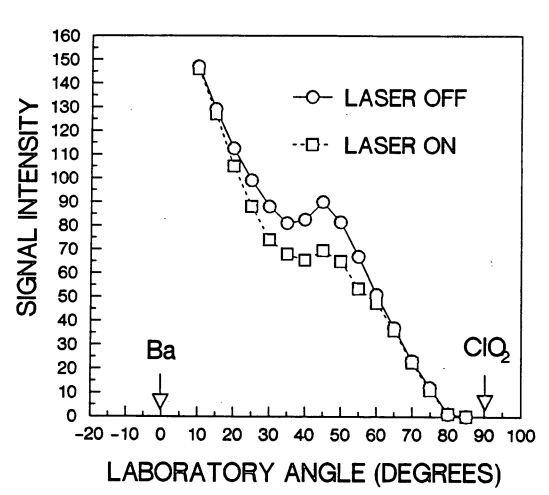


Fig. 3

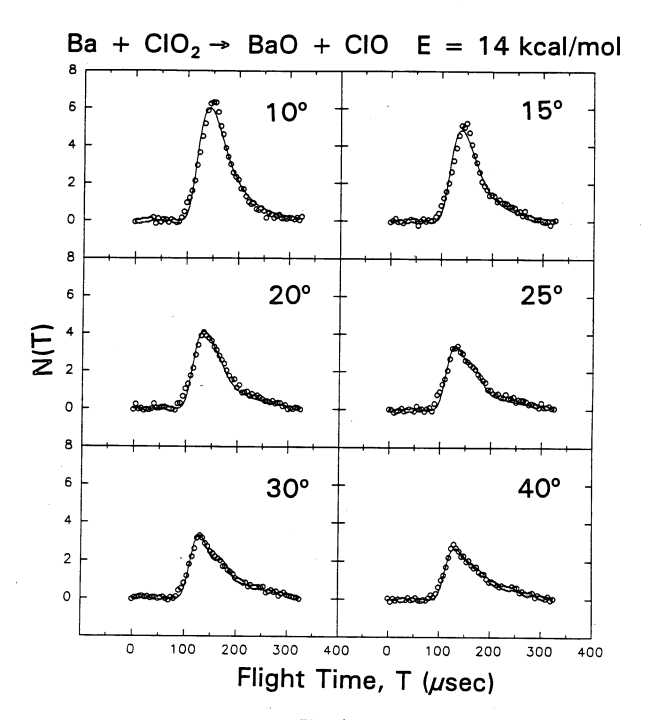
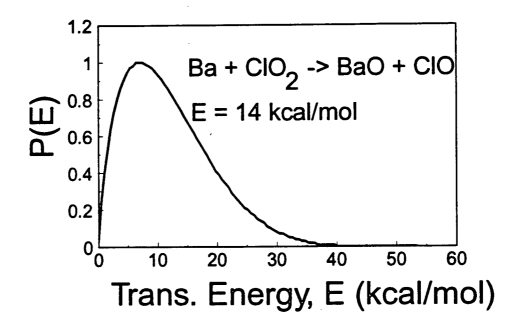


Fig. 4



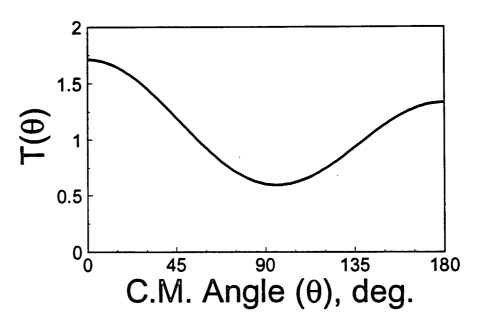
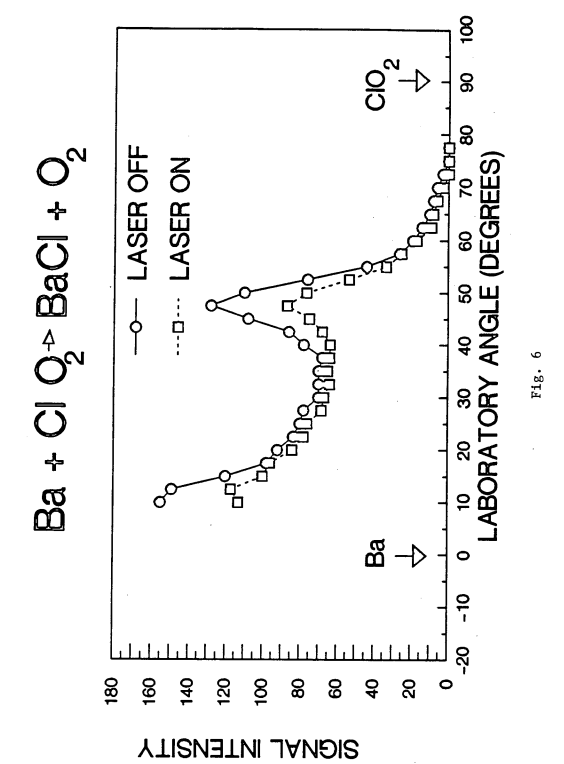


Fig. 5



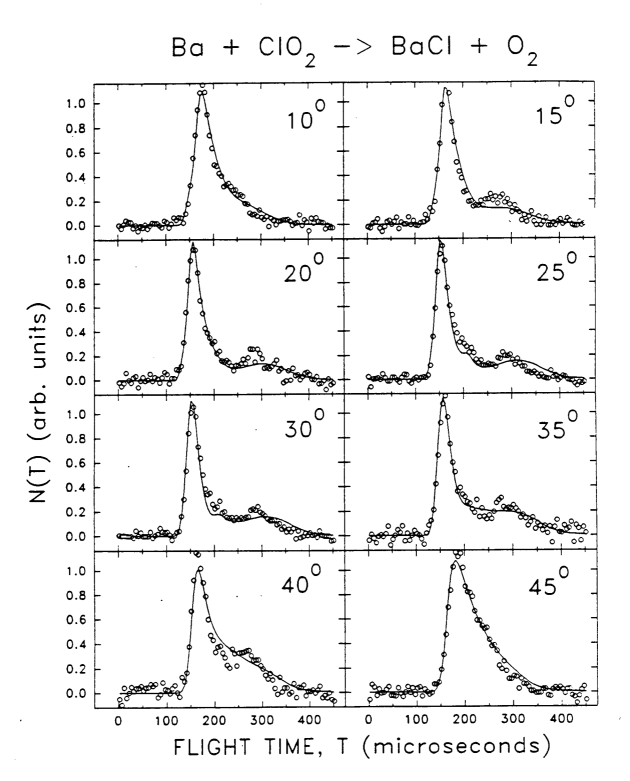
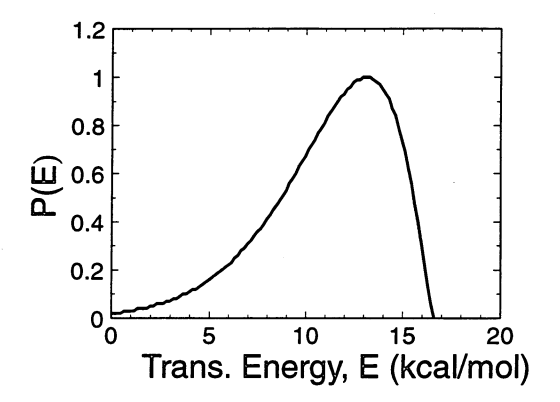


Fig. 7



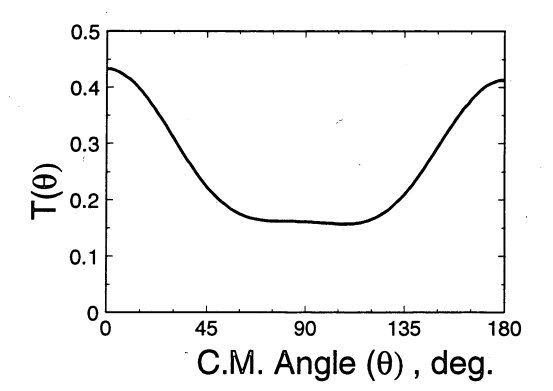


Fig. 8

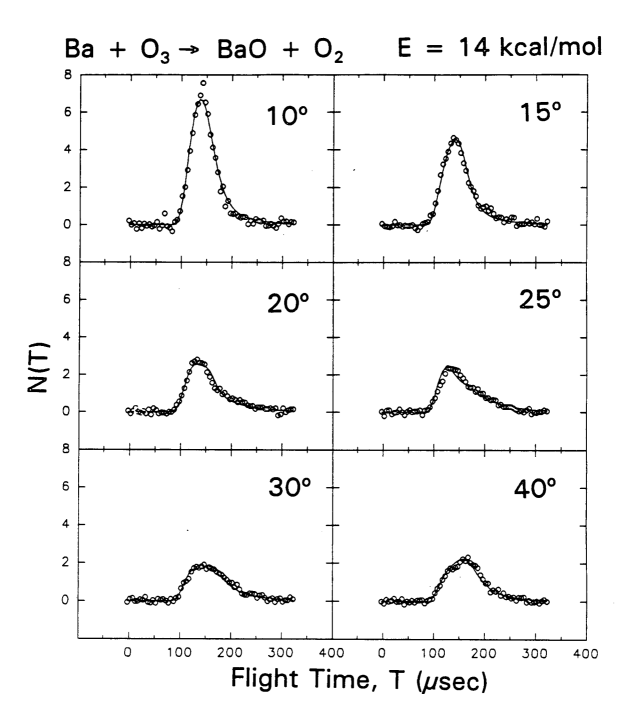
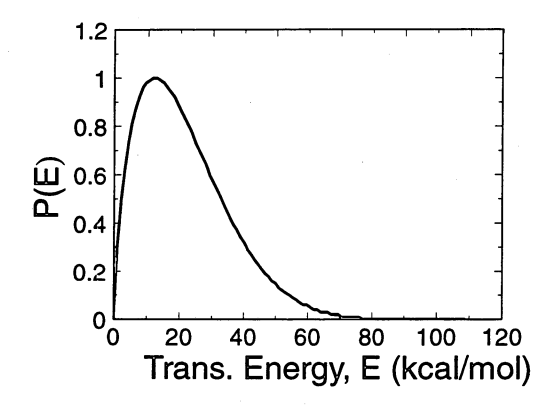


Fig. 9



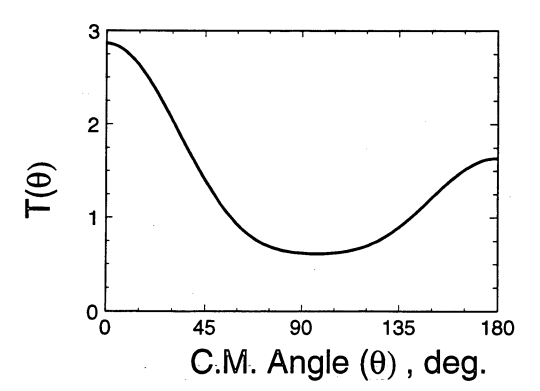


Fig. 10

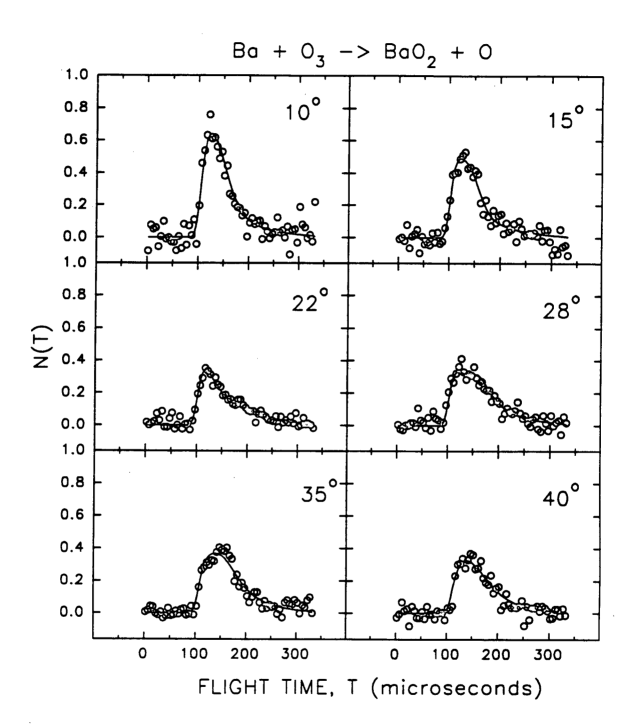
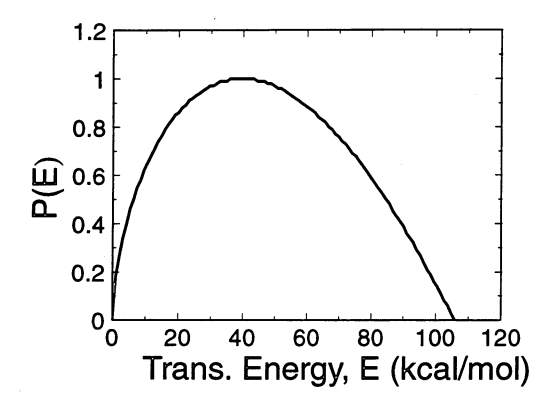


Fig. 11



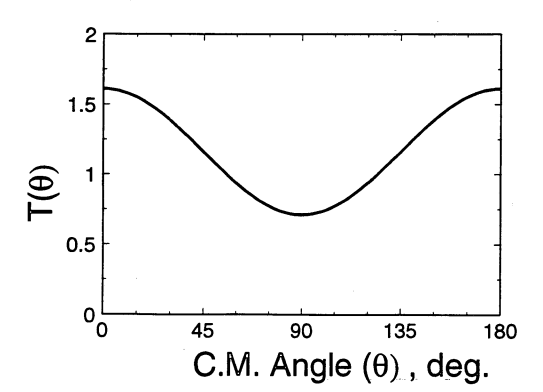


Fig. 12

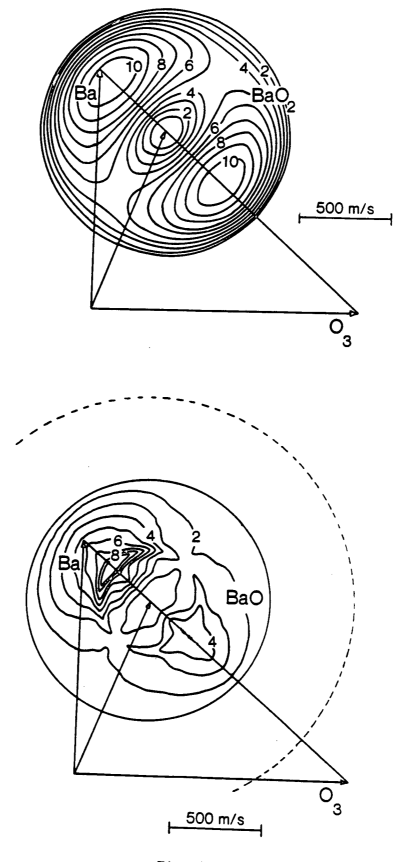


Fig. 13

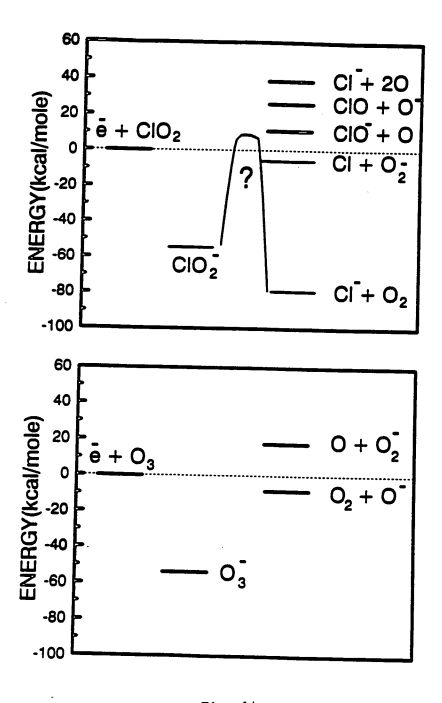


Fig. 14

CHAPTER 4

Photodissociation Dynamics of the NO₃ Radical.

ABSTRACT

The photodissociation of NO₃ was studied using the crossed laser-molecular beam technique at excitation wavelengths between 532nm and 662nm. Within this wavelength range, we observe formation of $NO_2(X^2A_1) + O(^3P)$ and $NO(^2\Pi) + O_2(^3\Sigma_p)$. is good evidence for substantial production of NO + $O_2(^1\Delta_g)$. The branching ratios for the two chemically distinct channels, NO₂ + O and NO + O₂ are strongly wavelength dependent. The observed yield of NO + O_2 reaches a maximum of $50\pm10\%$ near 590nm, falling to <0.02% below 584nm. Our observations indicate that strong vibronic coupling between the ²E' state and high lying vibrational levels of the ²A' state of NO₃ leads to formation of a vibrationally hot NO3 molecule that decays with kinetic competition between concerted unimolecular decomposition to NO + O2 and simple O-NO₂ bond rupture. Product time of flight spectra indicate that most of the excess energy in the molecular product channel is funneled into O₂ vibrational and electronic excitation, and NO + O₂ translational energy. From our experiments, we obtain a wavelength threshold for $NO_3(^2A_2') + h\nu \rightarrow NO_2(X^2A_1) + O(^3P)$ of 589 \pm 2nm, indicating that $D_o(O-NO_2) = 48.55 \pm 0.16$ kcal/mole. By combining this with the known values for $\Delta H_f(NO_2)$, and $\Delta H_f(O^3P)$, we calculate $\Delta H_f(NO_3) = 19.01 \pm 0.24$ kcal/mole (OK). Using heat capacity corrections, this corresponds to $\Delta H_1(NO_3) = 17.75 \pm 0.24$ kcal/mole (298K). We clearly observe structure in the NO + O₂ TOF at 588 and 594 nm due to vibrational excitation of the O₂ product. Our observations indicate that the potential energy barrier for $NO_3 \rightarrow NO + O_2$ ($^3\Sigma_g$, $^1\Delta$) is below $D_o(O-NO_2)$, i.e. <48.55 kcal/mol. Our results are compared to those of Magnotta and Johnston.

1. INTRODUCTION

Owing to its importance in the atmosphere¹, the nitrate free radical (NO₃) has received considerable attention experimentally¹⁻⁶ and theoretically.⁷⁻¹² Walsh¹³ predicted that NO₃ would have D_{3h} symmetry with a ground state configuration of ${}^{2}A_{2}$. Its excited electronic states are denoted by ${}^{2}E$ ' and ${}^{2}E$ ". The blue color observed upon mixing NO₂ with O₃ has been attributed to strong NO₃ visible absorption resulting from the ${}^{2}E$ ' \leftarrow ${}^{2}A_{2}$ ' transition.¹⁴ This absorption spectrum was first recorded in 1937 by Jones and Wulf,¹⁵ and reinvestigated by Ramsey¹⁴ in 1962, with several of the peaks assigned. The spectrum shown in Fig. 1 was recorded by Sander.¹⁶ The longest wavelength transition at 662nm was attributed by Ramsey¹² as due to the (0,0) band; the peaks at 623nm and 590nm result from a short progression in the symmetric stretching vibration, v_1 . Similar, relatively low resolution spectra have been recorded in a number of laboratories¹⁷⁻²⁰ using various chemical methods for generation of NO₃. These methods include:

Chemical Reaction:

a)
$$F + HNO_3 \rightarrow HF + NO_3$$
 (1a)

b)
$$NO_2 + O_3 \rightarrow NO_3 + O_2$$
 (1b)

Photochemical Decomposition:

$$ClONO_2 + h\nu \rightarrow Cl + NO_3$$
 (2)

(3)

Thermal Decomposition:

$$N_2O_5 \rightarrow NO_2 + NO_3$$

All evidence points towards the existence of a planar NO₃ molecule with a centrally located N atom. However, there has been considerable discussion³⁻¹² regarding the ground state symmetry of NO₃. It now appears that most inconsistencies which have arisen from assigning a rigid $C_{2v}(Y\text{-shaped})^3$ or D_{3h} (3-fold symmetric) structure, ^{6a} can be reconciled in terms of an NO₃ molecule possessing a triple minimum with only a small potential energy barrier for interconversion between three C_{2v} structures. ^{21,22} On the rotational timescale (10^{-12} s), molecular vibrations tend to average out the C_{2v} structure, and most features of NO₃ are consistent with an "average" D_{3h} structure. ^{21,22} If theory or experiment is sufficiently detailed, however, the underlying C_{2v} structure may become apparent. ^{21,22}

Sander observed a substantial temperature dependence for the NO₃ absorption cross section. ¹⁶ This has been supported by independent measurements by Ravishankara and Mauldin. ¹⁹ At nearly all wavelengths, the absorption cross section increases upon decreasing the cell temperature. ¹⁶ Such behavior suggests that the absorptions arise primarily from transitions from the ground vibrational state of NO₃, since the ground state population increases with decreasing temperature. Two peaks, at 678.4 and 637nm were found to show the opposite temperature dependence, ¹⁶ indicating that they are hot bands of the 662nm and 623nm transitions, respectively. Subsequently, Cantrell and coworkers²⁰ have conducted a more limited study at the 662nm (0,0) band only. They concluded that the 662nm absorption cross section is temperature *independent*. Such behavior, if real, could be taken as evidence for overlapping (0,0) and (1,1) transitions, possibly indicating a very small geometry change for the $^2E' \leftarrow ^2A_2'$ transition. The

nature of the NO₃ absorption spectrum, and its temperature dependence is of considerable importance in the present study and will be addressed in detail below.

The nitrate radical is one of the most poorly understood atmospheric molecules. Although the wavelengths of the peaks in the visible absorption spectrum are now well established, the identity of the transitions are only now becoming understood. However, there remains considerable uncertainty in the excited state dynamics of NO_3 . There is general agreement that in the absence of collisions, excitation at 662nm leads to fluorescence emission with a quantum yield of $1.0.^{3.58}$ Based on the JANAF values for $\Delta H_f(NO_3)$, $\Delta H_f(NO_2)$, and $\Delta H_f(O)$, as wavelength threshold of 583nm is calculated for the simplest possible photodissociation process in NO_3 :

$$NO_3(^2A_2') + h\nu \rightarrow NO_3(^2E') \rightarrow NO_2(X^2A_1) + O(^3P).$$
 (4)

This channel involves simple bond rupture, and no appreciable potential energy barrier is expected above the endothermicity.³ Consequently, this channel is expected to dominate at energies only slightly above threshold. Because of the substantial uncertainty $(\pm 5\text{kcal/mol})$ in the JANAF value for $\Delta H_f(NO_3, 0K)^{23}$ however, the wavelength threshold for process (4) is quite uncertain, lying between 528 nm and 649nm.

Although the concentration of NO_3 in the atmosphere has been found to be substantial at night, it is photolyzed rapidly at sunrise.⁵ If the quantum yield for reaction 4 is unity under the strong peaks in the absorption spectrum, then the formation and photolysis of NO_3 will lead to a null cycle in the balance of atmospheric O_3 :^{1,2}

$$NO_{2} + O_{3} \rightarrow NO_{3} + O_{2}$$

$$NO_{3} + h\nu \rightarrow NO_{2} + O$$

$$O + O_{2} + M \rightarrow O_{3} + M$$

$$Net: Null Cycle$$
(5)

However, several molecular channels are thermodynamically possible at all visible wavelengths: 1,2,23,25

$$NO_{3}(^{2}A_{2}') + h\nu \rightarrow NO_{3}(^{2}E') \rightarrow NO(^{2}\Pi) + O_{2}(^{3}\Sigma_{g})$$
 (6)
 $\rightarrow NO(^{2}\Pi) + O_{2}(^{1}\Delta_{g})$ (7)
 $\rightarrow NO(^{2}\Pi) + O_{2}(^{1}\Sigma_{g})$ (8)

The energetics of NO_3 photodissociation and an approximate correlation diagram, as postulated by Nelson, *et al.*³ are presented in Fig. 2. If the production of molecular $NO + O_2$ contributes significantly to the NO_3 integrated absorption cross section, then photodissociation of NO_3 in the atmosphere leads to catalytic destruction of O_3 by *visible* light:

$$NO + O_3 \rightarrow NO_2 + O_2$$

$$NO_2 + O_3 \rightarrow NO_3 + O_2$$

$$NO_3 + h\nu \rightarrow NO + O_2$$

$$2O_3 \rightarrow 3O_2$$
(9)

Net:

The only previous experimental investigations of NO_3 photochemistry were by Graham and Johnston¹ and later by Magnotta and Johnston.² Magnotta's study employed tunable laser flash photolysis of NO_3 generated by thermal decomposition of N_2O_5 in a

flow cell. Resonance fluorescence detection of the $O(^3P)$ and $NO(^2\Pi)$ photoproducts indicated the occurrence of both $NO_2 + O$ and $NO + O_2$ as primary photodissociation channels. Their results are summarized in Fig. 3. The quantum yield for $NO_2 + O$ was unity at 580nm, remained constant at 0.8 below 560nm and dropped to zero at 635nm. The $NO + O_2$ quantum yield peaked at 0.34 near 590nm, also dropped to zero to the red at 630nm, and fell to zero to the blue at 584nm.

A number of Magnotta's observations are particularly relevant to the present study. Formation of $NO_2 + O$ was observed at wavelengths considerably to the red of the nominal 583nm threshold implied by the JANAF values for $\Delta H_f(NO_3,0K)$. Although they did not attempt to explain this, more recent interpretations of their data by others³ suggest that it was collisionally induced. It is important to note that Magnotta's work was conducted *before* the first observation of fluorescence from NO_3 . Their measurements were "absolute"— knowledge of possible radiative processes was not required in the analysis. There was concern that the total measured photodissociation cross section $\Phi_{diss} = \phi_{NO} + \phi_O$ exceeded unity at 590nm,² indicating the presence of some systematic error. This is now believed to have resulted from minor heating of the flowcell by the resonance lamps used in detection of the photoproducts.²⁶ However, the mechanism of the dissociation, ie: direct or via internal conversion, collisional or collision free, was in question, as was the possible electronic state(s) of the O_2 photoproduct.

Recent work on the heat of formation of NO_3 has considerably complicated the interpretation of Magnotta's results. McDaniel⁴ and coworkers have reevaluated the heat of formation of N_2O_5 and NO_3 by vapor pressure and chemical equilibrium

measurements. Their value of $\Delta H_f(NO_3) = 15.39 \pm 0.75$ kcal/mol(298K) indicates that NO₃ is *more stable* than suggested by the JANAF value.²³ Combined with the well known values for heats of formation of $NO_2(X^2A_1)$ and $O(^3P)$,²³ these results lead to a calculated endothermicity of 51.17 kcal/mole for $NO_3 \rightarrow NO_2 + O(^3P)$, corresponding to a wavelength threshold of 554nm. This is even further to the blue of Magnotta's observed maximum in the NO_2 quantum yield.² If correct, McDaniel's value would indicate that a very large fraction of the photodissociation products observed by Magnotta and Johnston could not have resulted from photodissociation of ground state NO_3 under collision free conditions.

Quite recently, Weaver and Neumark²⁴ obtained photodetachment spectra of NO_3^- , and directly observed the "dark" ²E" state of NO_3 near 8000cm⁻¹. They also obtained a value of 3.937 ± 0.018 eV for the electron affinity of NO_3 . By combining this with the previously measured²⁷ value for $\Delta H_f(NO_3^-)$, they calculated $\Delta H_f(NO_3) = 17.9 \pm 0.8$ kcal/mol, in disagreement with McDaniel's result, but in good agreement with the less precise JANAF value of 17.0 ± 5 kcal/mol.

There are a number of possible explanations for Magnotta's observation of O atoms at wavelengths substantially longer than calculated using the JANAF tables,²³ or from the results of McDaniel⁴ or Weaver²⁴. The first possibility is that Magnotta's signal did result from photodissociation of a thermal population of NO₃ molecules, and all published heats of formation for NO₃ are simply far too low. This would have very important effects on calculated reaction rate constants for a number of atmospheric processes.⁴ If the stability of NO₃ was in fact much lower that previously believed, it

would also raise many questions in a number of systems involving NO₃, N₂O₅, and HNO₃. Another possibility is that Magnotta's O atom signal in the red resulted from two photon dissociation of NO₃:

$$NO_3(^2A_2)$$
 $\stackrel{hv}{\rightarrow}$ $NO_3(^2E')$ $\stackrel{hv}{\rightarrow}$ NO_3^{**} \rightarrow $NO_2^{\dagger} + O(^3P)$, (10)

In equation (10), NO₃** is an excited electronic state of NO₃ lying near 31,000cm⁻¹. The ionization potential of NO₃ is expected^{29,30} to exceed 80,000 cm⁻¹. Excited states of NO₃ are likely to exist at energies accessible by a two photon transition through the (²E') state. The NO₂ product could be vibrationally or electronically excited. Since the NO₃ fluorescence lifetime is known to be very long (30-300µsec),^{3,6a} a sequential two photon process through the ²E' state is quite possible, since a substantial population of NO₃* will accumulate during the laser pulse. If the second absorption step has a large cross section, it could easily be saturated. Two photon processes were identified by Magnotta² and explicitly taken into account in the analysis. However, if the second photon absorption was saturated, power dependence measurements could be misleading.

It is expected that a considerable fraction of the NO₃ molecules in the flowcell are vibrationally excited at room temperature. These species could be dissociated with light of lower energy than those in the ground state. The calculated vibrational populations for NO₃ at two temperatures are summarized in Table I. The large discrepancy between Magnotta's observed threshold (640nm) and that predicted from McDaniel's heat of formation (555nm) cannot be reconciled by arguing that internally hot molecules absorb

strongly. In Magnotta's experiment, it should be possible to characterize the NO₃ radicals by a Boltzmann distribution with a well defined temperature. Assuming McDaniel's value to be correct, an effective temperature of > 1000K would be required to provide sufficient internal excitation to lead to the observed wavelength dependence.

The participation of collisional effects, as suggested by Nelson, et al.³ is not easily dismissed. Magnotta's experiment was conducted at a pressure of several Torr. At this pressure, the excited NO₃ molecule could undergo many collisions within the fluorescence timescale. Our work was conducted under collision free conditions in a molecular beam. Thus, some additional insight into the possibility of collisional effects might be gained by comparison of Magnotta's observations with the present results.

Insight into the excited state dynamics of NO₃ may be obtained from past laser induced fluorescence (LIF) work.^{3,6a,21} Independently, and at nearly the same time, the groups of Nelson³, and Ishiwata^{6a} reported similar NO₃ fluorescence excitation spectra. These spectra were nearly identical to the NO₃ absorption spectrum (Fig. 1) at wavelengths above 615nm.³ This has been taken to indicate that the fluorescence quantum yield likely reaches 1.0 at a wavelength somewhere above 615nm.³ Upon tuning to the blue, however, the fluorescence intensity was found to decrease sharply, dropping to zero below 605nm.^{3,6a} This behavior clearly indicates the onset of efficient photodissociation below 605nm. Assuming that McDaniel's⁴ reported value of $\Delta H_f(NO_3)$ is correct, the LIF measurements would imply that the quantum yield for formation of NO + O₂ must be 1.0 in the wavelength range 555-605nm.^{2,3} This would conflict with Magnotta's results.²

Recently, the NO₃ system has attracted a considerable amount of theoretical work. 9-12,18 Although most studies have focussed on the question of the ground state geometry of NO₃, two groups have addressed the question of the height of the potential energy barrier between NO₃ and NO($^{2}\Pi$) + O₂ ($^{3}\Sigma$). The barrier height is of considerable importance in understanding the mechanism for oxidation of NO in the atmosphere, since it must involve an intermediate species, either NO₃ or N₂O₂.9 Using CASSCF and contracted CI methods, Siegbahn⁹ found that the C_{2v} constrained barrier height for $NO(^2\Pi) + O_2(^3\Sigma) \rightarrow NO_3$ was as high as 191 kcal/mole, and concluded that NO₃ was not an important intermediate in the atmospheric oxidation of NO to NO₂. Since the reaction NO + $O_2 \rightarrow NO_3$ is nearly thermoneutral (Fig. 3), Siegbahn's result implies that the reverse reaction will also have a very high potential energy barrier, at least when the reaction path maintains $C_{2\nu}$ symmetry. Subsequently, Boehm and Lohr²⁸ used ab initio methods to calculate the NO + O2 potential energy surface. Their results were also consistent with a very high potential energy barrier for the C_{2v} constrained reaction. They did find a considerably lower barrier for the NO + O₂ reaction when the path was nonplanar, and when the NO bond length was allowed to vary with O₂ bond distance.²⁸ However, in summarizing their work, it was stated that they would be surprised if it were determined that the transition state for NO + $O_2 \rightarrow NO_3$ had an energy outside 68 ± 17kcal/mole.²⁸ Their lowest limit (51kcal/mole) would imply a wavelength threshold for NO₃ \rightarrow NO($^2\Pi$) + O₂($^3\Sigma$) of 560nm. If the LIF^{3,6a} and NO₃ enthalpy data⁴ is correct, and the quantum yield for $NO_3 \rightarrow NO + O_2$ is 1.0 for 555nm $< \lambda < 605$ nm, the calculations might suggest that the O_2 is formed exclusively in an electronically excited state.

Many observations regarding NO₃ indicate that strong vibronic coupling exists between the first optically allowed ²E' state and high lying levels of the ground ²A' state. ^{21,24}. This coupling, also known as the Douglas effect³⁰ is well known in NO₂ and SO₂, and leads to a number of important properties. In the case of NO₃, the absorption spectra are found to be diffuse, ^{14,31} even at relatively high resolution, at wavelengths where dissociation cannot occur. This results from a very high level density, on the order of 1000/cm⁻¹, ^{3,32} leading to a very congested spectrum. This results in a very long fluorescence lifetime of 30-300μsec.³ In the "zero order" picture, this can be explained in terms of strong "dilution" of the initially prepared state due to strong vibronic coupling to nearby high lying levels of the ground state. More precisely, the wavefunctions describing the NO₃ eigenstates accessed by absorption of a visible photon are superpositions of the ²A' and ²E' states. This effect will be shown to be of considerable importance in the present work.

Most of the uncertainties in NO₃ largely stem from its reactivity, the resulting need for complex reaction mixtures for its generation, and the possible role of collisional processes under bulk analysis. A number of these complicating factors have been eliminated in the present study. The results of Magnotta,² combined with the questions regarding $\Delta H_f(NO_3)$,^{4,23,24} prompted this experimental effort. We have used the crossed laser-molecular beam method to study the photodissociation of NO₃ under "isolated" collision free conditions. Our goal was to identify the chemical products and their electronic state(s) and to determine barrier heights and possible mechanisms for the

dissociation as a function of wavelength. Here we report the angular and velocity distributions for both product channels resulting from excitation of NO_3 between 532 and 662nm. Based on the NO_2 + O translational energy distributions, and the wavelength threshold for this product channel, we conclude that NO_3 is substantially *less stable* than is implied by the recent thermochemical measurements.⁴ Our result is in excellent agreement, but somewhat more precise, than the recent calculated value based on EA_{NO3}^{24} and $\Delta H_f(NO_3^-)$.²⁷ This system has proven to be rather complex. Consequently, the project has evolved over a period of time, with the necessity of several experimental approaches. Some of the early conclusions from this work were presented in the thesis of Bongsoo Kim.³³ As will be shown, although many conclusions were correct, the early work did suffer from several shortcomings. As a result of our recent experimental efforts, the deficiencies in our earlier work have been identified, and a clear picture regarding the photodissociation dynamics of NO_3 has emerged.

II. EXPERIMENTAL

The original experiments were conducted using the A-machine, which is the same apparatus used for the Ba atom experiments described in Chapters 1-3. The experimental arrangement for photodissociation experiments is shown in Fig 4. The most recent experimental results were obtained using a second apparatus specifically designed for studies of photodissociation dynamics.³⁵ Unlike the A-machine, it has a fixed detector and rotatable molecular beam source (Fig. 5). This "Rotating Source Machine" (RSM) is capable of achieving considerably better translational energy resolution. This is due

to a longer neutral flight length (36.7cm vs. 20.8cm), but similar effective ionizer length (~1cm). An additional feature of the RSM is that it is possible to extend the neutral flight length to ~85cm, providing exceptionally high resolution. Of course, this is only possible if the signal levels are high enough to withstand a decrease in signal intensity by a factor of ~5. Perhaps the only disadvantage of the RSM is a more constricted source chamber, requiring a more compact molecular beam source, and a more limited pumping capacity.

A continuous NO₃ radical beam was formed by pyrolysis of a 1-5% N₂O₅/He mixture in the nozzle tip at ~ 600 K. Nitrogen pentoxide (N₂O₅) was prepared by reaction of NO₂ with O₃ as described by Schott and Davidson.³⁶ The N₂O₅/He mixture was formed by passing He through a large glass trap containing crystalline N₂O₅, held at \sim -20C in a thermostatically controlled bath. The trap was identical to the Si gel traps normally used to trap O₃ (Chapter 3). In the present application, the Si gel was removed. A number of nozzle designs were used in this study. The first nozzle (Nozzle 1-Fig. 6a) was made by drilling a 0.020" dia. hole in a 0.020" thick plate, 0.25" in diameter, and welding it to the end of a 0.25" OD, 0.15" ID stainless steel tube. Calculations, 33 summarized in Fig. 7, indicated that high NO₃ concentrations (>0.1 Torr before expansion) may be achieved, but only using pyrolysis times shorter than 1 msec at temperatures near 600K. By heating only the final 2mm tip of the nozzle, the heating time for the N₂O₅ molecules could be kept short. The newly formed NO₃ radicals are rotationally cooled by supersonic expansion into the vacuum chamber before being destroyed by thermal reactions which would normally lead to formation of NO and NO2.

The heater was fabricated by soldering to the end of the nozzle a small copper sleeve (0.25"ID, 0.35"OD, 0.15" long); wrapped and soldered to the sleeve was one turn of Thermocoax wire heated by DC current (10A, 5V). A 2" long water cooled copper block was soldered to the tube 1mm behind the heating zone to keep the remainder of the nozzle at room temperature. A slot was machined into the circumference of the nozzle tube to decrease the heat loss from the hot nozzle tip to the water cooled region (Fig. 6)

In the early stages of this project, the only known method for monitoring the NO₃ concentration was by its LIF intensity at 662nm.^{3,6a,33} The nozzle temperature, total stagnation pressure, (typically 150 Torr), and N₂O₅ concentration were adjusted to maximize the NO₃ fluorescence. Once these conditions were found, the beam was sampled directly by the mass spectrometer through a 0.003" dia. pinhole. It was found that the parent N₂O₅ molecule yielded essentially no NO₃⁺ fragment upon 200V electron impact ionization. However, the NO₃ molecule resulting from its thermal dissociation showed a strong parent ion peak. This remarkable behavior allowed us to optimize the nozzle conditions by simply maximizing the m/e=62 (NO₃⁺) countrate while monitoring the beam directly. This was found to be extremely useful since NO₃ does not fluoresce at most wavelengths used in this work.^{3,6a} Based on the magnitude of the photodissociation signal that we observe, we estimate the NO₃ concentration in the nozzle before expansion to be in the range 0.1 to 0.5 Torr. This is extremely high for an unstable radical and made it possible to obtain very high signal to noise ratios in the photofragment time-of-flight spectra.

In this experiment, beam collimation was especially important. This is because the beam is actually dominated³³ by NO₂. Other species, namely NO, O₂, and N₂O₃ are also present to a lesser extent (Fig. 7). All of these impurities will contribute to a large, constant background at product masses. Only NO₃ and NO₂ absorb at the wavelengths studied.^{1,2,6a} However, NO₂ remains bound upon single photon absorption.²³ Consequently, these contaminant molecules cannot contribute to the photodissociation signal because they are constrained to the beam. The detector only observes those laser-correlated particles scattered out of the beam due to photodissociation. In order to gauge the possible role of two photon absorption by NO₂, parallel experiments were run using 5% NO₂/He mixtures. Formation of NO + O was observed at 532nm only when the laser was tightly focussed and very high pulse energies (>50mJ/pulse) were used. Multiphoton effects in NO₂ could not be seen at longer wavelengths. In the experiments reported here, the laser was unfocussed with a 2mm spot size at the interaction region, and the pulse energies were in the range 0.5-15mJ/pulse.

In early experiments using Nozzle 1, we estimated the NO₃ vibrational temperature.³³ This involved measurements of hotband intensities at 678.4nm relative to the (0,0) band at 662nm. This hotband corresponds to excitation of NO₃ molecules having 360cm^{-1} of vibrational excitation. The hotband intensity ratios measured by Sander, et. al.,¹⁶ at two well defined temperatures were employed in the calculation of the beam temperature. Our measurements indicated that essentially no vibrational cooling results from supersonic expansion through the 600K nozzle with 0.020" channel length (i.e. $T_{vib} \approx 600\text{K}$).³³ This is consistent with the general expectation that vibrational

cooling in supersonic expansions is relatively inefficient.

As can be seen in table I, due to the very low vibrational frequencies of NO₃, even at 600K a very significant fraction of the molecules in the beam are vibrationally excited. This is of considerable importance in our determination of the O-NO₂ bond dissociation energy since internally hot molecules will be dissociated with redder light than those in the ground state. For a given laser wavelength above threshold, these hot molecules could lead to photofragments with larger translational energy than would be found from dissociation of ground state molecules. In an effort to cool vibrations, the front plate and nozzle orifice was cooled to ~300K by spot welding six nickel wires, 0.030" dia, 1" long to the plate surface, without constricting the orifice. The wires were soldered to a water cooled 0.125" dia. copper tube. This arrangement did result in some vibrational cooling, based on the observation of decreased hotband intensities, as described elsewhere.³³ However, the use of Ni wires as described is not a very efficient, nor elegant way to provide a heat sink. In the final experimental effort on the RSM, a somewhat more sophisticated nozzle was designed (Fig 6b). One advantage of the newest design was that it provided an even shorter pyrolysis time. This is clearly desirable since calculations indicate that this will increase the NO₃ concentration³³ (Fig7). Of course, the chief goal of this design was to reduce the vibrational temperature of the NO₃. As can be seen in Fig. 6, the region around the nozzle orifice is cooled by the water cooled front electrode to a temperature near 300K. In principle, this should result in some cooling of the NO₃ radicals during expansion through the nozzle. This expectation is based on the earlier measurements using the older nozzle designs in which the hot band intensity near 675nm was probed using LIF, and found to decrease if the aperture was cooled.³³ In general, we found that the vibrational temperatures of the particles in the beam are reasonably close to the ambient temperatures of the nozzle orifice. To date, quantitative LIF measurements have not been conducted using the present nozzle. However, we are able to distinguish between signal resulting from photodissociation of ground state and vibrationally hot NO₃, as will be shown conclusively in the discussion section.

Nozzle 2 (Fig 6B) was heated by passing AC current (1V, 60A) from the front electrode to the back electrode. Both electrodes are copper, and are water cooled. The 0.25" OD copper tubes used for the cooling water also facilitated transmission of the electric current. The front wall of the stainless steel nozzle tube was turned down to 0.010" thickness. The flow of current through this region preferentially heats this thin walled section of the tube. The length of this region was 2mm. The end of the tube was capped, and the nozzle orifice (0.020" dia.) centrally located. Soldered to the front of the tube, near the orifice, was the water cooled front electrode. Due to the large heat capacity of this electrode and efficient cooling due to the flow of water, the region near the nozzle orifice is expected to be near 300K. Since the heated portion of the tube was very thin, it had little mechanical strength. Consequently, the front electrode is actually supported by a ceramic insulating sleeve. This provided strength, while at the same time provided a "slip fit" to allow for expansion of the thin walled portion of the tube upon heating. We have found that this nozzle is very reliable and provides a stable NO₃ beam with reproducible velocity and intensity.

The velocity of the molecular beam was determined by modulating the molecular beam with a spinning slotted disk in front of the detector. The beam velocity could also be checked without using the wheel, in holeburning experiments. Due to the large absorption cross section of NO₃, ^{1,2,16} at high laser powers it was relatively easy to observe depletion of NO₃ parent molecules in the beam by simply monitoring the NO₃⁺ parent ion signal while sampling the beam directly through the 0.003" aperture. The time profile of this depletion could be used to determine the beam velocity. This was particularly convenient in the early experiments using the A-machine since the wheel had to be removed from the apparatus during the photodissociation experiments.

A number of different laser systems have been employed in this project. For most experiments, a XeCl excimer pumped Lambda Physik FL2002 pulsed dye laser was used to excite the NO₃ radicals. The laser polarization could be rotated using a double fresnel rhomb. In the earliest work, a Quanta-Ray Yag pumped pulsed dye laser was employed. Although this laser was capable of achieving very high pulse energies, the repetition rate was only 20Hz. Owing to the very large NO₃ absorption cross section, the laser power had to be attenuated to minimize multiphoton effects. Thus, the excimer pumped laser, which could be operated at 50 Hz, but with a considerably lower pulse energy, was preferable. A series of experiments were also conducted on the RSM using a continuous wave Coherent 699 ring dye laser pumped by a 6W Ar⁺ laser. The dye laser was operated in "broadband" mode (ie. like a 699-05), and the typical output was ~1 Watt. The laser beam was modulated using the cross correlation technique³⁷ by sending the beam through a small vacuum chamber in which a double sequence cross

correlation wheel was spun at 392 Hz. The cross correlation wheel provided 50% duty cycle with a nominal 5μ sec time resolution. In all CW experiments, the laser wavelength was monitored by sending a very small portion of the beam through a 0.5m monochromator and the transmission was measured using a visible PMT. The monochromator was calibrated using a standard neon resonance lamp.

In the RSM, a small fraction of the photodissociation products fly 36.7 cm into an electron bombardment ionizer.³⁵ The distance on the A-machine is only 20.8cm. However, the detectors are similar in both machines.^{34,35} The resulting ions are extracted into a quadrupole mass filter and detected by a Daly ion detector. The geometry of the detector defining slits ensures that only the 2x2mm zone centered at the interaction region is viewed by the detector. Consequently, only those dissociation events which occur within $\sim 2\mu s$ are detected. A 1024 channel scaler with $1\mu s$ dwelltime, triggered by the laser pulse, was used for product TOF measurements. In all experiments, the detector was operated at unit mass resolution.

The translational energy distribution P(E) for each product channel is obtained by the forward convolution method.³⁵ The experimental time of flight data is compared to a calculated spectrum based on a trial P(E). The trial P(E) is iteratively adjusted until good agreement is observed between the calculated and observed TOF data. The computer program used in this procedure convolutes over the spread in beam velocity and the known detector parameters, and calculates the required laboratory to center of mass Jacobian transformations needed for the simulation.

III. RESULTS AND ANALYSIS

In most experiments, the laser was vertically polarized, perpendicular to the detector axis. However, if the dissociation lifetime is less than one parent rotational period, an anisotropic product angular distribution is possible, and must be considered in the data analysis. We have found that the product angular distributions are isotropic at all wavelengths studied—i.e. the anisotropy parameter (β -parameter) is zero. This measurement was accomplished by conducting a series of polarization experiments at a number of wavelengths down to 532nm. Using a reduced laser intensity to minimize possible saturation effects, product time-of-flight spectra were recorded using a number of polarizations. The polarization was rotated using a double Fresnel rhomb. The isotropic product angular distribution that we observe is consistent with a dissociation timescale considerably longer than one NO₃ rotational period (~1psec).

Experiments were conducted at a wide range of laser wavelengths, primarily at the peak NO₃ absorbances in the range 532-662nm. Photodissociation product signal was observed at m/e=46 (NO₂⁺), $m/e=32(O_2^+)$, $m/e=30(NO^+)$, and $m/e=16(O^+)$. The earliest work obtained using the A-machine and nozzle 1 will be discussed in section 3.1. The purpose of this section is to present a number of the important observations. In the earliest work, we focused our attention on the peak near 590nm, as photolysis at this wavelength led to the maximum observed NO + O₂ signal level. The effect of tuning the laser above and below 590nm (still using nozzle 1 on the A-machine) is explored in detail in section 3.2. We clearly observed multiphoton effects at wavelengths above 590nm. Although this effect could be minimized by running experiments at very low

power, the signal to noise ratio was very poor when the laser output was reduced to these low levels. This observation laid the groundwork for the more recent experiments, which were conducted using the RSM. In addition to eliminating a substantial contribution from dissociation of hot NO_3 radicals, the RSM experiments provided much higher resolution, making it possible to resolve vibrational structure in the $NO + O_2$ TOF. This data, obtained using the excimer pumped dye laser will be presented in section 3.3. The focus of section 3.4 is on measurements pertaining to the determination of $D_o(O-NO_2)$. This quantity was obtained from two separate kinds of measurements. The first method involves observation of the wavelength dependence of the $NO_2 + O$ channels. This data was obtained under single photon conditions using the ring laser. The second method involved careful measurements of the maximum translational energy release for the $NO_2 + O$ channel at wavelengths considerably above threshold. These two complimentary measurements, as well as a number of other observations, lead to an accurate determination of $D_o(O-NO_2)$, allowing us to calculate $\Delta H_f(NO_3)$ directly.

III. A. Photodissociation of NO₃ at 590nm using A-machine and Nozzle 1.

The Newton diagram for the NO channels from photodissociation of NO_3 at 590nm is shown in Figure 8. The radius of the Newton circle for the NO_2 product from the NO_2 + O channel will be very sensitive to the actual value of $D_o(O-NO_2)$ and the vibrational energy of the NO_3 molecules in the beam. The radius must be small, however, because of the close proximity to the wavelength threshold for NO_2 + O. In addition, the heavy NO_2 particle is constrained to appear near the beam because of its

light O atom recoil partner. The simultaneous existence of the NO₂ + O and NO + O₂ product channels at 590nm is demonstrated in the 590nm data shown in Fig 9. The observed NO₂⁺ signal results from the NO₂ + O channel. A similar slow peak is also seen in the NO⁺ TOF. This is the daughter ion from fragmentation of NO₂ to NO⁺ upon electron impact ionization. The fast peak in the NO+ TOF is due to NO parent ion from the $NO + O_2$ channel. This conclusion is confirmed by the existence of a similar peak in the m/e=32 (O_2^+) TOF spectrum corresponding to the momentum matched O_2 fragment (Fig 9). Signal from both dissociation channels was found to disappear if the nozzle heater was switched off. It also decreases upon heating above 600K due to decrease in NO₃ concentration. In the early experiments, we did not have any means to rotate the laser polarization without realignment of the laser beam. However, based on the comparable signal levels in experiments using vertical and horizontal laser polarization, we believed the effect was near zero, and this was confirmed in more recent work using the RSM. The NO TOF spectra at other detector angles are shown in Figs. 10 and 11. An additional feature, observed between 10 and 20 degrees at $\sim 100-170 \mu s$ results from sequential 2 photon absorption by NO₃ yielding NO₂ + O (Fig 10). This contribution to the observed TOF was not fitted in the analysis. The two photon effect will be discussed below. As expected from the kinematics, the NO₂ product channel is only observed at angles less than or equal to 12.5 degrees, whereas NO + O₂ products can be seen at all detector angles. (Figs. 10,11)

The product translational energy distributions that resulted from the fit of the 590nm data (solid line in Figs. 10,11) are shown in Fig 12. Very little structure could

be resolved in this data. However, upon closely examining the NO + O_2 TOF, a small but reproducible shoulder can be observed on the fast edge of the NO time of flight spectra at all angles. This fast shoulder must result from formation of $NO(^2\Pi) + O_2(^3\Sigma_g)$ (the ground electronic states of O_2). This conclusion is based on energy conservation-product translational energies higher than 25 kcal/mol leave insufficient energy for excitation to the $O_2(^1\Delta_g)$ state. Interestingly, the dominant, slower component in the TOF spectrum fell completely within the translational energy range dictated (by conservation of energy) for the NO + $O_2(^1\Delta_g)$ channel. These observations prompted the use of two separate contributions in fitting the NO + O_2 translational energy distributions. The two components were fitted by requiring that the slower peak fall completely within the energy boundaries consistent with NO + $O_2(^1\Delta_g)$ (i.e.; $E_{trans,max} \sim 25$ kcal/mol). These early observations strongly suggested that $^1\Delta_g$ is in fact the dominant electronic state of O_2 resulting from the photodissociation of NO_3 at 590nm.

The branching ratio for the $NO_2 + O$ to the $NO + O_2$ channel at 590nm, based on measurements using the relatively hot NO_3 radicals formed in nozzle 1, was obtained from the m/e = 30 (NO⁺) TOF data. The optimized fit of the entire data set (Figs. 9-12) acquired at all detector angles yields an "apparent" branching ratio R_{app} defined by equation 1:

$$R_{app} = \frac{\chi (NO + O_2)}{\chi (NO_2 + O)} = 1.10$$
 (11)

where the χ 's are the integrated areas under the product P(E) distributions for the two different chemical product channels, normalized for the differing Jacobian factors calculated by the computer program in carrying out the LAB \rightarrow CM coordinate transformations. The actual branching ratio R is related to the "apparent" ratio by the following expression:

$$R = R_{app} \times \frac{\sigma_{ion}(NO_2)}{\sigma_{ion}(NO)} \times \frac{F(NO^+/NO_2)}{F(NO^+/NO)}$$
(12)

In equation 12, $\sigma_{ion}(NO_2)$ and $\sigma_{ion}(NO)$ are the ionization cross sections for NO_2 and NO_3 , respectively. They were calculated using the following empirical relationship³⁸ relating the peak ionization cross section (σ_{ion}) to molecular polarizability (α):

$$\sigma_{ion} = 36\sqrt{\alpha} - 18 \tag{13}$$

The molecular polarizabilities are taken to be the sum of the atomic polarizabilities.³⁹ $F(NO^+/NO_2)$ is the fraction of NO_2 product molecules that fragment to NO^+ , a quantity directly measurable in this experiment. Although the fragmentation pattern of NO_2 is expected to strongly depend on its level of internal excitation, our fragmentation ratios agree quite closely to published values for room temperature NO_2 .⁴⁰ This similarity is a result of our close proximity to the wavelength threshold for $NO_3 \rightarrow NO_2 + O$ at 590nm. By inserting the appropriate values into equation 2 we find that R=1.01,

indicating that yields of the two chemical products are equal at 590nm to within the errors associated with the calculation, estimated to be \pm 20%. The largest source of error stems from uncertainty in the P(E) for E_{trans} < 1kcal/mol due to our inability to detect photodissociation products within 5 degrees of the beam. Thus we conclude that at 590nm:

$$NO_3 \rightarrow NO_2 + O \Phi_1 = 0.5 \pm 0.1$$
 (14)

$$\rightarrow$$
 NO + O₂ $\Phi_2 = 0.5 \pm 0.1$ (15)

This calculation is based on several assumptions. First, we assume that photodissociation occurs within 2μ sec; i.e., before the excited NO₃ molecules fly out of the viewing range of the detector. We have also assumed that $\Phi_1+\Phi_2=1.0$ at 590nm. This appears to be a reasonable assumption based on the absence of fluorescence under collision free conditions below 600nm. We also assume that multiphoton processes are negligible at 590nm. Although this will be discussed in detail below, this appears to be be reasonable.

III. B. Wavelength Dependence Studies on the A-machine

A dramatic decrease in the NO + O_2 quantum yield is observed upon tuning the laser from 590 to 584nm as evidenced by the NO TOF data recorded at 10 degrees (Fig 13). As expected, the same behavior is observed in the O_2 ⁺ data. The integrated signal

intensity for the NO₂ + O channel (measured at m/e = 30 or 46) was found to increase upon tuning from 590 to 584nm. Figure 14 shows the NO⁺ time of flight data taken at 10 degrees as the photolysis wavelength is set above 590nm. The laser pulse energy was 10mJ/pulse. Two major effects may be noted. First, the yield of NO + O_2 (faster peak) decreases somewhat with respect of the NO₂ + O peak. This decrease in NO + O₂ was also apparent from the O₂⁺ TOF data. The second observation is that another component grows in, between the two peaks. On reaching 596nm, the three components have nearly merged together. We attribute this third component, which was also observed at NO₂⁺ (but not O₂⁺), as sequential two photon absorption of NO₃ resulting in formation of NO₂ + O. All available evidence indicates that two photon absorption does not lead to formation of NO + O₂. Because the two photon process leads to NO₂ with substantially larger translational energy than the one photon process, we at first believed that they could be distinguished from one another. However, it is impossible to rule out the possibility of formation of internally excited NO₂ from two photon photodissociation of NO₃. This makes it very difficult to unambiguously separate the two contributions, particularly since the signal to noise ratio declines considerably at lower pulse energies. However, it is certain that the contribution from two photon effects increases substantially upon tuning the laser above 590nm, even at the lowest laser power. It should be noted that power dependence measurements showed linear behavior, suggesting that the second photon absorption cross section is very large, and was saturated at these laser powers.

Prolonged averaging at m/e=32 (O_2^+) with the laser at 605nm revealed weak

signal indicating the existence of the NO + O₂ channel. No O₂+ signal was observed at 613, 623, 637, or 662nm (local maxima in the room temperature NO₃ absorption spectrum), but the NO₂ + O channel could be seen with decreasing intensity at wavelengths up to 637nm, using the pulsed laser. It should be noted that it was impossible to rule out possible contributions from some multiphoton effects in the long wavelength regime. There was also considerable concern about the role of vibrationally hot molecules, which might be dissociated considerably to the red of true NO₂ + O wavelength threshold. Figure 15 shows the observed wavelength dependence for the two chemically distinct channels, measured using Nozzle 1. Because the absorption spectrum is not known for rotationally cold NO₃, the data in Fig. 15 is not normalized for the absorption cross sections. While the data in Fig. 15 was reproducible, and was obtained under the lowest possible laser fluence (<1mJ/pulse), there was concern that the NO₂ + O signal at wavelengths above 590nm could still result from multiphoton effects or from photodissociation of internally hot NO₃. Power dependence studies showed that the intensity was linear in power at these pulse energies, but this could also be due to saturation of the second photon transition. It is interesting to note that the wavelength dependence for the NO + O₂ signal level (Fig. 15) shows many similarities to that obtained by Magnotta (Fig. 3). In particular, the sharp drop of the $NO \, + \, O_2$ channel below 590nm that we observe is essentially identical to their earlier result. Our quantum yield of 0.5 ± 0.1 at 590nm is slightly higher, but in reasonable agreement with Magnotta's value. The main difference is that the NO₂ yield in our experiment is smaller than that obtained in Magnotta's experiment in the red region of the spectrum. The question of the NO₂ quantum yields near threshold will be addressed in section 3.4.2.

III. C. High resolution studies on the RSM using Nozzle 2.

As already mentioned, the longer neutral flightlength of the RSM provides the opportunity to achieve increased resolution in the translational energy distributions. Structure observable in the NO + O_2 data obtained on the A-machine suggested that it might in fact be possible to resolve features resulting from vibrational excitation of the product molecules. Resolving such structure could provide useful insight into the dynamics of the NO + O_2 channel, particularly if this structure is wavelength dependent. A series of experiments were conducted at two excitation wavelengths, 588nm and 594nm. These wavelengths were chosen because the NO + O_2 channel is appreciable, yet they fall near the edges of the range where NO + O_2 is observed (Fig 15). Earlier experiments suggested that the structure in the TOF changed somewhat, upon tuning across this wavelength region.

The NO⁺ fragment was monitored at a number of angles for each wavelength. For each time of flight, the data was accumulated for a period of 1 hour. The data is shown in Figs. 16 and 17, and the resulting translational energy distributions are shown if Figs. 18 and 19. A considerable amount of structure is observed in the time-of-flight spectra obtained at high resolution. The resulting translational energy distributions reflect this structure. The P's(E) shown in Figs. 18 and 19 result from the best fit to the data at all angles for each wavelength.

III. D. NO₂ + O Translational energy distributions at wavelengths above threshold

An accurate determination of the enthalpy of formation of the nitrate free radical (NO₃) is of considerable importance in properly modelling reactions involving nitrogen oxides and ozone. The currently accepted literature values^{4,23} of $\Delta H^o_f(NO_3)$ rely on the combined results of several separate thermochemical measurements involving dinitrogen pentoxide (N₂O₅), a relatively unstable molecule that is extremely difficult to prepare in pure form.

In addition to providing considerable information on the dynamics of the photodissociation process, this technique is a particularly attractive method for thermochemical measurements involving unstable species.³⁵ In the photodissociation of an isolated NO₃ molecule, conservation of energy dictates that:

$$E_{photon} + E_{int, NO_3} = D_o(O-NO_2) + E_{trans, NO_2+O} + E_{int, NO_2} + E_{int_{iO}}$$
 (16)

In a simple bond rupture, such as $NO_3 \rightarrow NO_2 + O(^3P)$, the maximum translational energy release corresponds to the production of ground state fragments. Thus, by measuring the maximum translational energy for the $NO_2 + O$ products, we obtain $D_o(O-NO_2)$ directly:

$$E_{photon} + E_{int,NO_3} = D_o(O-NO_2) + E_{trans,max,NO_2+O}(17)$$

or

$$D_o(O-NO_2) - E_{int,NO_3} = E_{photon} - E_{trans,\max,NO_2+O}(18)$$

Since the photon energy is known, measurement of $E_{trans, max, NO2 + 0}$ leads directly to $D_o(O-NO_2)$ - $E_{int,NO3}$. Usually, E_{int} is assumed to be zero. However, in the present case, this could be a bad assumption, particularly because the molecules are formed by pyrolysis. It is also important to note that because NO_3 has low vibrational frequencies, a substantial fraction of the beam will be vibrationally hot (Table I). Consequently, in the present experiment we have conducted measurements at several wavelengths. Based on the maximum translational energies at different excitation wavelengths, we should be able to address the possible role of vibrationally excited NO_3 in the beam.

In the earlier experiments on the A-machine, we focussed our attention primarily on wavelengths above 580nm since this is the wavelength region where the NO + O_2 channel was seen. It also seemed reasonable to measure the maximum translational energy for the NO_2 + O channel near threshold, since the v^2/u^2 Jacobian factor in the CM-LAB transformation leads to large signal levels for slow particles. This appeared to be a logical approach because we were very concerned with the contribution from multiphoton processes. However, the maximum translational energies that we obtained at 590nm were near 4kcal/mole (Fig. 12). The majority of the signal, however fell at energies below 2kcal/mole. Unfortunately, it was impossible to tell for certain whether the fast edge resulted from hot molecules in the beam, from multiphoton effects, or from dissociation of "cold" NO_3 .

In planning the final experimental effort on the RSM, we considered the possibility of multiphoton effects and photodissociation of hot molecules in the molecular beam. It appears that multiphoton effects can be largely attributable to the long

dissociation timescale for NO_3 at $\lambda > 590$ nm. Thus, it appeared that the best approach on the RSM would be to measure the maximum translational energy release at a number of wavelengths considerably above threshold, where the dissociation timescale is expected to be much faster. Since it is clear that the quantum yield for $NO_2 + O$ is 1.0 below 580nm, any structure in the translational energy distribution observed at higher excitation energies might reflect the vibrational distribution of the parent NO_3 molecules. A "cold" nozzle (Fig. 6B) was designed in hopes of cooling the radicals—if a substantially different result was seen using this nozzle, we would have evidence for dissociation of hot NO_3 .

The TOF data taken at m/e=16 (O⁺) at an excitation wavelength of 560nm is shown in Fig. 20. Three translational energy distributions are shown in Fig. 20. As shown in Fig. 20b, the best fit was obtained with the maximum recoil energy of 3.5 kcal/mol. Note that most products have an energy of ≤ 2.5 kcal/mole. Figures 20a and 20c show the sensitivity of the fit. It is clear that $E_{trans, max} = 3.5 \pm 0.5$ kcal/mol at 560nm. Figures 21 and 22 show the TOF and P(E) for the NO₂ + O channel at 548nm. The maximum translational energy release was 4.3 kcal/mole at 548nm. The fast edge of the translational energy distributions at both wavelengths were assumed to correspond to formation of NO₂ in the ground vibrational and rotational state. This assumption is reasonable in view of the proximity to the wavelength threshold and the nature of the process: simple O-NO₂ bond rupture. The calibration of the laser is known to be within 5cm^{-1} (0.014kcal/mol) based on the experiments described in Chapter 5.

Based on these measurements, we obtain the following relationships:

548nm:

$$D_o(O-NO_2) - E_{int,NO3} = E_{br} - E_{trans, max} = (52.15 - 4.3) = 47.8 \pm 0.5 \text{kcal/mol}$$
 (19)

560nm:

$$D_o(O-NO_2) - E_{int,NO3} = E_{hr} - E_{trans, max} = (51.03 - 3.5) = 47.5 \pm 0.5 \text{ kcal/mol}$$
 (20)

We are encouraged by the fact that the maximum recoil energy follows the laser photon energy.

If all of the NO₃ molecules in the beam were in their ground vibrational state, then the maximum product translational energies would give an upper limit to the bond dissociation energy. Because the beam was formed at 600K, however, low frequency vibrational modes of NO₃ are expected to have appreciable populations. We must consider possible contributions from these hot molecules. The vibrational frequencies ¹⁰ for NO₃ (assuming D_{3h} symmetry) and expected equilibrium populations relative to the ground vibrational state are listed in Table 1. Based on the earlier measurements of hotband intensities, we have found that the vibrational temperature of the NO₃ beam is very close to the nozzle temperature. Using the first nozzle, we obtained a temperature of 600K, in close agreement with the expected nozzle temperature. In nozzle 2, the orifice is water cooled, and is expected to be nearer to room temperature. Unfortunately, it is not clear that the vibrational temperature of the NO₃ will be as cool as this. However, we can confidently exclude the possibility that the NO₃ is hotter than 600K since it is formed by thermal dissociation of N₂O₅. Since the NO₃ molecules are formed

by an endothermic process (simple bond rupture), it is extremely unlikely that they will be hotter than the ambient nozzle temperature, particularly since they subsequently undergo collisions before and during the supersonic expansion.

The temperature dependence of the NO₃ absorption spectrum (Fig. 1) indicates that the transitions at 548 and 560nm do not result exclusively from hotband activity. Although Cantrell²⁰ has found the absorption at 662 nm to be temperature *independent*, there is no evidence that the earlier temperature dependence measurements of Sander¹⁶ suffer from wavelength dependent artifacts. Thus, the molecules absorbing at 560 and 548nm exhibit behavior similar to that seen across nearly the entire spectrum. It is also worth noting that the observation that the maximum translational energy release tracks with the photon energy gives us confidence that we have not accidentally selected 'hotband' transitions at these two wavelengths.

III. E. Wavelength dependence for the $NO_2 + O$ Channel

A second approach exists for determining the O-NO₂ bond dissociation energy. It involves measuring the NO₂ + O yields as a function of wavelength near threshold. Based on our earlier experiments on the A-machine, and the results presented in section 3.4.1, it appeared that the wavelength threshold for formation of O + NO₂ was near 600nm. As already mentioned, very strong multiphoton effects are seen in this region, as discussed in Section 3.1.

The pulsed dye laser is typically operated at 50Hz, and the pulse duration is ~ 20 ns. This translates into a duty cycle of $1x10^{-6}$ -- i.e. the laser is only on for one-

millionth of the time. With a total output power of up to 1W, it is not surprising that multiphoton effects are observed in the NO₃ photodissociation experiments at wavelengths where the excited state lifetime likely exceeds many picoseconds. The Coherent 699 ring dye laser provides comparable output power (1W), but it is continuous, rather than pulsed. Using the cross correlation technique, we can obtain a 50% duty cycle and still obtain product TOF information. With the ring laser, the effective peak power is 106 times lower than in the pulsed experiment, ruling out the possibility of multiphoton effects.

The Newton diagram for the O atom product from photodissociation of NO₃ at various energies above threshold is shown in Fig. 23. By monitoring the light O atom fragment at 10 degrees, we can observe the NO₂ + O reaction channel provided that the photon energy is at least 0.25kcal/mol above threshold. The O⁺ time of flight spectrum obtained at an excitation wavelength of 592nm using the ring laser is shown in Fig. 24. The data was accumulated for a period of 1 hour. Due to the inherently stable output of the ring laser, it was very easy to maintain a constant output power for many hours throughout the duration of the experiment. The integrated area under the O⁺ TOF peak is plotted as a function of wavelength in Fig. 25. The signal was undetectable at wavelengths above 599nm. The signal level increased steadily at wavelengths below 599nm, reaching a plateau near 592nm. The signal level increased sharply below 584nm. Similar experiments were conducted while monitoring NO from the NO + O₂ channel at 40 degrees. The shape of the TOF was found to be very similar to that obtained using the pulsed laser, indicating that multiphoton processes do *not* contribute

significantly to the observed NO + O_2 TOF (Fig. 26). Of course, it multiphoton activity is competing with the NO + O_2 yield, this will lead to an artificially low value for the NO + O_2 quantum yield. The integrated intensity for NO at 40 degrees is plotted as a function of wavelength in Fig. 27. It is interesting to note that the signal drops to zero below 586nm, in excellent agreement with the behavior seen earlier by us³³ using the pulsed laser, and by Magnotta.³

Based on the O⁺ wavelength dependence plotted in Fig. 25, The "observed" wavelength threshold for the NO₂ + O channel was 599 ± 2 nm (E_{hr}= 47.71 ± 0.16 kcal/mol) under single photon conditions using the ring laser. However, it is important to note that at an angle of 10 degrees, the minimum recoil energy detectable is 0.25 kcal/mol. Thus the threshold for formation of NO₂ + O with zero recoil energy is $47.71\pm0.25 - 0.25 = 47.46 \pm 0.16$ kcal/mol. Assuming that at least some of the NO₂ products are formed in their ground rotational and vibrational states near threshold (a good assumption for simple bond rupture), we obtain the following relationship:

$$D_o(O-NO_2) - E_{int,NO3} = E_{thresh} = 47.46 \pm 0.16 \text{ kcal/mol}$$
 (21)

This value is slightly lower, but in reasonable agreement with the values obtained from the maximum translational energy release at energies well above threshold.

In order to establish a value for $D_o(O-NO_2)$, we must evaluate $E_{int,NO3}$. Based on the data presented in Table 1, it is clear that a large fraction of NO_3 molecules will possess one quantum of excitation in the doubly degenerate v_4 mode at 363cm⁻¹.

Depending on the actual temperature of the beam, this population could even be 84% of that having v_4 =0. The population of NO₃ molecules having two quanta of doubly degenerate v_4 excitation (2 x 363cm⁻¹=726cm⁻¹) will not be negligible if the beam temperature is near 600K. It is important to note that 363cm⁻¹ translates to 1.03kcal/mole, and 726cm⁻¹ = 2.07 kcal/mole. Thus, the potential error in our value of $D_o(O-NO_2)$ will be large if we do not correctly evaluate $E_{int, NO3}$. Further discussion will be deferred to section 4.1.

IV. DISCUSSION

A. NO₃ Photodissociation Wavelength Dependence.

Perhaps the most striking result of this work is the narrow wavelength range over which NO + O_2 is observed, corresponding to a range of less than 1.75 kcal/mole of excitation energy. In contrast, NO₂ + O is observed with a quantum yield of 1.0 at all wavelengths below 584nm. At 590nm, NO₃ is excited only 5.3 kcal/mole above the 2 E' band origin. Formation of NO + O_2 involves substantial bond rearrangement requiring bond distortion far greater than that provided by only 5.3 kcal/mole of vibrational excitation. As suggested by Nelson,³ this, together with the strong vibronic coupling known to exist^{21,22,24} between the 2 E' excited state and high lying levels of the 2 A' ground state, strongly suggests that formation of NO + O_2 involves internal conversion of NO₃. Siegbahn has calculated a barrier height of 145 kcal/mole for the C_{2v} constrained dissociation process. Our results clearly indicate that the barrier actually lies below 48.11 kcal/mole ($\lambda > 594$ nm) above NO₃. This could suggest that the symmetry of NO₃

may not be near C_{2v} at the transition state, since the potential barriers for transition states of lower symmetry are expected to be substantially less than 145 kcal/mole.²⁸ We will address this point in detail in discussing the NO + O₂ TOF spectra in section 4.3. It is worthwhile noting that our experimental barrier height is slightly below the lowest limit of 51kcal/mol calculated by Boehm and Lohr.²⁸

The sharply decreasing signal level for $NO_3 \rightarrow NO + O_2$ below 590nm is a very dramatic effect. Our observations are in very good agreement with those of Magnotta.² Since this falloff was reproduced using both the pulsed and ring laser, it cannot result from participation of multiphoton effects. Using the first nozzle, we found that the quantum yield for the channel decreases from 0.5 ± 0.1 to <0.01 over a 6nm range, corresponding to a change in photon energy of only 0.50 kcal/mol. In order to explain this behavior, we must also consider the NO₂ + O signal level in this wavelength interval. The only reliable $NO_2 + O$ data above 590nm is that obtained using the ring laser under strict one photon conditions. We find that the $NO_2 + O$ yield increases rapidly at photon energies above the 599nm threshold, reaching a plateau near 590nm. This behavior is quite different than that seen earlier using the pulsed laser. The large difference indicates that a substantial fraction of the NO₂ signal we saw earlier at wavelengths above 590nm resulted from multiphoton effects. Although we expect that the NO₃ will be cooler in the present experiment due to the cooling of the nozzle orifice, based on Table I, the large shift in the NO₂ + O threshold cannot be simply attributed to a cooler NO_3 beam. We also observe a very sharp increase in observed NO_2 + O signal level (at 10 degrees) at a wavelength below 584nm. Since no other processes are possible in this wavelength range, our observations indicate that kinetic competition is occurring between the two photodissociation channels.

According to eq. 21, $D_o(O-NO_2) - E_{int.NO3} = 47.46 \pm 0.16$ kcal/mol. We might be tempted to postulate that the energy threshold for dissociation of ground state NO₃ is 47.46 kcal/mol, i.e. $E_{int,NO3} = 0$ for absorption near 599 nm. However, this conclusion is incorrect. This "threshold" in fact corresponds to dissociation of NO₃ radicals which posses 1 quantum of v_4 excitation at 363 cm⁻¹(1.03kcal/mol). Based on this, we calculate that the NO₃ bond dissociation energy is $D_o(O-NO_2) = 47.46 + 1.03 = 48.49$ kcal/mol. This corresponds to a wavelength threshold of ~ 589 nm. A 589nm threshold for dissociation of ground state NO₃ to NO₂ + O explains the sharp decrease in NO + O₂ yield seen in our experiments and in the experiments of Magnotta. Upon tuning the laser through the NO₂ + O threshold at 589nm, that channel will quickly dominate since the rate of simple bond rupture, even near threshold, will be much greater than that for concerted formation of NO + O_2 . In Fig. 25, a sharp increase in m/e=16 (O⁺) signal was seen below 584nm. This sharp increase results from dissociation of ground state NO₃ to NO₂ + O. It must be remembered, however, that we are able to detect the O⁺ product at 10 degrees only if E_{trans} ≥ 0.25 kcal/mol. By drawing a smooth line through the data to the wavelength axis, the threshold for this process is found to be near 585nm $(E_{h\nu}=48.85~kcal/mol)$. The energy threshold for formation of "zero kinetic energy" NO_2 + O is 48.85 - 0.25 = 48.6 kcal/mol. This corresponds to a wavelength of 588nm, which is 1nm to the blue of the 589nm threshold stated above. However, the slight deviation from 589nm is not unreasonable since an extrapolation is required in a region where there is an overlapping contribution from dissociation of vibrationally hot NO₃.

All of our observations using the ring laser provide evidence that the wavelength threshold for dissociation of ground state NO₃ to NO₂ + O is 589 \pm 2nm and that D_o (O-NO₂) = 48.55 \pm 0.16 kcal/mol. However, these measurements are based on threshold behavior, and will be incorrect if any appreciable potential energy barrier exists for NO₃ \rightarrow NO₂ + O. Calculations of D_o(O-NO₂) from the translational energy distributions well above threshold at 560 and 548nm do not suffer from this possible source of error. Based on our 560nm data, we calculated that D_o(O-NO₂) - E_{int, NO3} = 47.5 \pm 0.5 kcal/mol. This is in excellent agreement with the threshold measurements if the fast component in the P(E) (Fig. 20b) results from photodissociation of NO₃ with one quantum of 363 cm⁻¹ excitation, i.e. E_{int,NO3} = 1.03 kcal/mole. At 548nm, we found that D_o(O-NO₂) - E_{int, NO3} = 47.8 \pm 0.5 kcal/mol. This is also in good agreement with the 589nm threshold, provided that E_{int,NO3} is 1.03 kcal/mole.

It is perhaps ironic that by measuring the NO + O_2 quantum yield at 590nm, we have in fact simply measured the population of NO₃ in the ground vibrational state. At 590nm, the photon energy is just below threshold for the NO₂ + O channel. Since fluorescence is known to be zero, our measurement that ϕ_{NO+O2} = 0.5 ± 0.1 at 590nm simply reflects the ground state population, all of which is dissociating to NO + O₂. Of course, the other half of the NO₃ molecules in the beam are internally excited, and are dissociated to NO₂ + O, since E_{photon} + $E_{int,NO3}$ exceeds $D_o(O-NO_2)$. Because the potential energy barrier for NO + O₂ is just below the endothermicity for NO₂ + O, the concerted NO + O₂ channel occurs readily, but only within a very narrow wavelength

range.

It is rather straightforward to show that the 589 ± 2 nm "threshold" corresponds to the threshold for dissociation of ground state NO₃. If the threshold for dissociation of ground state NO₃ was actually at 599 nm, it would be very difficult to rationalize a number of our observations. First, we know that there must be internally hot NO₃ molecules in the beam. This conclusion is based on our earlier hotband intensity measurements³³ and the calculated data in Table 1. If we attributed the observed threshold for $NO_2 + O$ at 599nm to be due to ground state NO_3 , we imply that internally excited NO₃ molecules make no contribution here. However, we have found that the 599nm "threshold" and the maximum translational energy release at 560nm and 548nm are mutually self- consistent. This would lead us to the conclusion that the absorptions at 560 and 548 also do not result from contributions from hot NO₁. This would be surprising since vibrationally hot molecules must contribute to the NO₃ absorption spectrum, and the temperature dependence at 560 and 548 is similar to that seen at nearly all other wavelengths. It also seems very unlikely that absorption at both 560 and 548 have no contribution from vibrationally excited NO₃ since the photon energies for these two transitions lie 390 cm⁻¹ apart. If the 548nm transition resulted from excitation from the ground state, then that at 560nm should show some contribution from excitation of those NO₃ molecules having one quantum of 363cm⁻¹ excitation to the same upper state. Based on these arguments, it appears to be nearly impossible that hot molecules can make no contribution at 560, 548, and 600nm. Secondly, this scenario would imply that the $NO + O_2$ and the $NO_2 + O$ channel both have wavelength thresholds near 600nm, and quantum yields rising in tandem to ~ 0.5 at 590nm. Such behavior could not arise from simple kinetic competition between the two channels, since the Arrhenius preexponential factor for the simple bond rupture will be much greater than for the NO + O₂ channel. Thus, if both channels had the same wavelength threshold, the NO₂ + O channel should have been dominant at 590nm.

It might seem possible that the 599nm threshold corresponds to threshold for dissociation of NO₃ molecules possessing two quanta of 363cm^{-1} vibrational excitation, i.e. $E_{int} = 2.07$ kcal/mol. In this case, we would calculate a value for the O-NO₂ bond energy of 47.46+2.07 = 49.53kcal/mol. This corresponds to a wavelength threshold of 577nm. This is clearly impossible since we find that the quantum yield for NO₃ \rightarrow NO₂ + O has already reached ~ 1.0 at wavelengths 7nm to the red at 584nm.

Using the ring laser, the NO + O_2 channel showed the same behavior as was seen earlier using the pulsed laser. This indicates that multiphoton effects do not have a significant effect on the NO + O_2 wavelength dependence. However, multiphoton effects can effect the NO + O_2 quantum yield. For example, at 594nm, we predict that a ground state NO₃ molecule will dissociate exclusively to NO + O_2 . However, if the excited NO₃ absorbs a second photon it will dissociate to NO₂ + O. Thus, multiphoton effects could artificially lead to a lower observed NO + O_2 quantum yield.

In earlier pulsed experiments, although we did see some O_2 at 605nm, it was very weak. In Magnotta's experiment, however, $NO + O_2$ was observed with a significant quantum yield up to 620nm, with a small tail out to 630nm. Since we obtained very similar wavelength behavior using both the pulsed laser and the dye laser, it appears that

sequential absorption by NO₃ does not lead to formation of NO + O₂. There are several possible explanations for the differences between our NO + O₂ yield above 595nm and We do not expect Magnotta's NO₃ vibrational temperatures to be that of Magnotta. much different than ours. Thus, it is very unlikely that the difference can be attributed to a larger contribution from vibrationally hot NO₃ in Magnotta's experiment. However, due to efficient rotational cooling in the beam, we do expect a considerable difference in rotational temperature in the two experiments. In our experiment, a rotational temperature well below 50K would be expected, whereas that in the flow cell should be near room temperature. The rotational constants of NO₃ are on the order of 1cm⁻¹. At room temperature, there will be a substantial population of NO_3 with J=20-30. These rotational levels correspond to rotational energies of approximately 1.5-3 kcal/mole, which at this energy is in fact a 15-30nm wavelength range. Thus, it appears that a considerable fraction of the NO + O_2 seen by Magnotta to the red of 600nm (Fig. 28) could arise from dissociation of rotationally hot NO₃. Unfortunately, since the dependence of the NO₃ absorption spectrum on rotational temperature is not known, it is difficult to be more quantitative in addressing this possibility. Of course, collisional effects could also play a role in the NO + O₂ signal at wavelengths above 605nm in Magnotta's experiment.

Our experiment is only sensitive to those dissociation processes which occur within $\sim 2\mu$ sec of the laser pulse. Under normal circumstances, this is not a significant consideration since the dissociation timescales of even very large molecules seldomly exceeds several hundred picoseconds. However, in the present case, it is possible that

the $NO_3 \rightarrow NO + O_2$ dissociation timescale could become very long above 600nm. If it is on the order of microseconds, the NO_3 molecules will "fly out" of the viewing region before dissociation. We attempted to observe this in the present experiment using two approaches. First, we attempted to monitor NO_3 depletion in the beam with the detector at 0 degrees using the ring laser. We also attempted experiments with the laser moved upstream of the interaction region. These experiments have been inconclusive to date. Although a microsecond dissociation timescale might seem surprising for a 4 atom molecule, it is in fact quite reasonable in view of the fact that the onset of fluorescence emission is seen at 605nm and the fluorescence lifetime is known to exceed 30μ secs at 613nm. However, since fluorescence is seen at 605nm, it appears that the wavelength range in question is very small. This conclusion is also supported by RRKM calculations which indicate that for a 4-atom molecule, the range of internal energies corresponding to microsecond $NO + O_2$ dissociation timescales will be very limited.

Perhaps the most important conclusion regarding the observed wavelength dependence of the two photodissociation channels is that the measured quantum yields will be very sensitive to the vibrational population of the NO_3 molecules. Based on our results, we predict that the $NO + O_2$ quantum yield for a pure sample of "cold" NO_3 would be unity at 590nm. A real sample of NO_3 at finite temperature will have some vibrationally excited molecules, and so the $NO + O_2$ yield will always be smaller than this. Multiphoton effects will also serve to diminish the observed $NO + O_2$ quantum yields at wavelengths above 590nm.

IV. B. Calculation of $\Delta H^{o}(NO_3)$

All of our experimental data indicates that the wavelength threshold for O-NO₂ bond rupture of ground state NO₃ is 589 ± 2 nm. Based on the dramatic decrease in NO + O₂ below 590nm, it is nearly impossible to rationalize a value below the energy corresponding to 589nm. In the case of NO₂, it is known^{41,42} that the fluorescence quantum yield drops sharply to zero near 397.92nm. Apparently, the quantum yield for NO + O becomes substantial within only a few cm⁻¹ of the dissociation threshold. In the case of NO₃, the NO₂ + O channel should also become dominant near threshold, in the absence of a significant potential energy barrier in the exit channel.

It should be noted that our translational energy distributions at 560nm and 548nm peak away from zero. This might be taken as evidence for a potential energy barrier in the exit channel, invalidating our measurements of the threshold wavelength for NO₂ + O. However, it is extremely unlikely that the observed P(E) indicates the existence of an exit barrier based on the fact that we did get reasonable agreement in the threshold measurement and the measurements of E_{trans,max} above threshold. Thus one might wonder why the P(E) peaks away from zero when we are so close to threshold for a reaction without a barrier. The most probable answer is that at energies this close to threshold, the range of accessible product internal energies is not "continuous" on our energy scale since, of course, the NO₂ vibrational energy is quantized. Only a few vibrational levels of NO₂ are available at these energies. Consequently, the peaking away from zero likely reflects the small number of NO₂ quantum states available in this energy range. For a given NO₂ vibrational level, excess energy can only be channeled into relative

translational motion or NO_2 rotation. Thus, even though the excitation energy will be rapidly randomized in NO_3 due to strong vibronic coupling effects, the P(E) peaks away from zero. This suggests that a very limited amount of energy appears in rotation. This is not at all surprising for simple O- NO_2 bond rupture since the impact parameter for the ensuing O + NO_2 "half collision" will be near zero due to the D_{3h} symmetry of NO_3 . In the photodissociation of NO_2 , on the other hand, the O + NO_3 impact parameter is large and a substantial fraction of the available energy can appear as rotational motion in the NO_3 fragment.

We have concluded that the O-NO₂ wavelength threshold for ground state NO₃ is 589 ± 2 nm. This indicates that $D_o(O-NO_2) = 48.55 \pm 0.16$ kcal/mol. This result is more precise than the values we obtained from measuring the maximum translational energy release at 560 and 548nm. However, our value of 589 ± 2 nm as the threshold for $O + NO_2$ is solidly supported by our observed wavelength behavior using both the pulsed and ring laser. It is also supported by the earlier results of Magnotta. In order to calculate $\Delta H^o_f(NO_3)$, OK, we require values for $\Delta H^o_f(O^3P)$, OK and $\Delta H^o_f(NO_2)$, OK. According to the JANAF tables, $\Delta H^o_f(O^3P) = 59.98 \pm 0.02$ kcal/mol OK and $\Delta H^o_f(NO_2) = 8.59 \pm 0.19$ kcal/mol OK. The uncertainty in the latter quantity will considerably increase the uncertainty in our calculated value of $\Delta H^o_f(NO_3)$, OK.

In a beautiful series of experiments, Robra and Welge⁴¹ studied the photodissociation of NO₂ near threshold in a molecular beam. They monitored NO₂ fluorescence and NO photodissociation product intensity simultaneously while scanning their photolysis laser through the NO₂ dissociation threshold. They found that

fluorescence dropped to near zero abruptly near 397.93nm. Based on their observations, they concluded that the energy threshold for dissociation of ground state NO₂ to NO + O is 25130.6 \pm 0.6cm⁻¹. This corresponds to a value for D_o(O-NO) of 71.8604 \pm 0.0002kcal/mol. Based on the JANAF values for $\Delta H^o_f(O^3P) = 58.98 \pm 0.02$ kcal/mol (0K) and $\Delta H^o_f(NO) = 21.46 \pm 0.04$ kcal/mol (0K), we calculate $\Delta H^o_f(NO_2) = 8.58 \pm 0.06$ kcal/mol (0K). This is nearly identical to the JANAF value, but the error limits are considerably smaller. Based on this, we can use our value for D_o(O-NO₂) to calculate $\Delta H^o_f(NO_3) = 19.01 \pm 0.24$ kcal/mole at 0K. Based on published heat capacity corrections⁴³ which agree with those we calculate according to Herzberg,⁴⁴ our value corresponds to a room temperature value of $\Delta H^o_f(NO_3) = 17.75 \pm 0.24$ kcal/mole (298K).

The value for $\Delta H^o_f(NO_3)$, 298K obtained in this experiment is in excellent agreement with the calculated value employing the NO_3 electron affinity of Weaver²⁴ and the previously measured value of $\Delta_t H^o(NO_3)$. They calculated $\Delta_t H^o(NO_3) = 17.9 \pm 0.8$ kcal/mole (298K). However, our finding is not consistent with the thermochemical measurements of Cantrell.⁴ Their value for $\Delta_t H^o(NO_3)$ would imply that the threshold for formation of $NO_2 + O$ from ground state NO_3 lies near 556nm. If this were the case, we should have observed the onset of a significant low energy component in the 548nm product time of flight spectra due to the onset of dissociation of ground state molecules. This was not observed. Moreover, since NO_3 fluorescence is known to drop to zero below 605nm, their result could only indicate that the quantum yield for $NO + O_2$ is unity for $560 < \lambda < 605$ nm, which is ruled out by this work, and that of Magnotta.

The enthalpy of formation that we have obtained is 2.36 kcal/mol higher than the recent thermochemical values. Unfortunately, these values, which are clearly incorrect, have been incorporated by JANAF in the most recent thermochemical tables.⁴³ The consequences of using our results, rather than those newly adopted, will be very substantial in certain cases. The effect will be particularly dramatic in calculations pertaining to processes that are nearly thermoneutral. For example, by using our value, the equilibrium constant for $NO_3 + HCl \rightleftharpoons HNO_3 + Cl$ is calculated to be approximately 1000 times larger than the value based on Cantrell's thermodynamic data.⁴

It is important to point out that based on Magnotta's earlier results, and our observation of $NO_2 + O$ considerably to the red of 600nm in the pulsed experiments, we believed for a long time³³ that the true threshold was above 600nm, and that even the value obtained by Weaver²⁴ for $\Delta H^{\circ}_{f}(NO_3)$ was near the lower edge of our error bar. It was only after completing the "single photon" experiments using the ring laser that we realized that multiphoton effects were in fact dominating our measurements of the $NO_2 + O$ channel above 590nm. We had been somewhat "deceived" by the earlier experiments. It appears unlikely that the vibrational energy of the NO_3 observed in the flowcell could be much hotter than that we derive from thermal decomposition of N_2O_5 . However, in Magnotta's experiment, we do expect considerable contribution from photodissociation of higher rotational levels from the thermal distribution of NO_3 molecules, and this will increase the contribution to the $NO_2 + O$ signal to the red of 589nm. In retrospect, it is perhaps not too surprising that NO_3 does suffer from multiphoton activity above 590nm, since the groundstate molecules are below the O- NO_5

bond energy and are likely to be dissociating on an extremely long timescale. Our work indicates that an electronic state of NO₃ exists near 31000 cm⁻¹, and this state dissociates readily to NO₂ + O, with the NO₂ internally excited. It should be noted that 31000cm⁻¹ is an energy range considerably above that currently known for NO₃. In addition, because the selection rules for one and two photon absorption processes are different, it is very likely that this state will not be accessed by one 320nm photon.

IV. C. The Structure in the NO + O_2 TOF.

Our high resolution spectra show considerable vibrational structure. Since we have established that the NO signal results primarily from photodissociation of NO₃ in its ground state, the observed structure must result from internal excitation of the product molecules. This reaction channel involves formation of two diatomics, each with vibrational and rotational degrees of freedom. In view of this, it is somewhat surprising that this amount of structure is observed at all. Intuitively, we would expect that NO₃ \rightarrow NO + O₂ will lead to a significant yield of vibrationally excited O₂, since no O-O bond exists in NO₃, but a very strong O=O bond is produced in the reaction. This hypothesis is clearly supported by the data. We first examine the 588nm data. As summarized in Table II, the substantially larger vibrational frequency for NO allows us to rule out the possibility that the structure in the P(E) is attributable to a progression in NO vibrations. The spacing that we observe clearly corresponds to progressions in O2 vibrational motion. Figure 29 shows the expected maximum translational energy release for various internal states of the O₂ product if we assume that the NO partner is internally cold (v=0). Our value for $\Delta H_1(NO_3)$ was used in the calculations. The O_2 vibrational

energies compiled in Huber and Herzberg⁴⁵ were also employed, as this appears to be the most reliable data on O2 available. Of course, our detector is not state specific, and we must be very careful when drawing conclusions about the internal states of the products. It is worthwhile mentioning that if we had higher resolution, we would likely be able to resolve more structure in the high translational energy range. The TOF becomes very compressed at energies above 25 kcal/mol and possible structure in the TOF is not resolvable under the present conditions. However, since we do see significant NO + O₂ with translational energy above 25 kcal/mole, we are able to state with certainty that at least some $O_2(^3\Sigma)$ is formed in the dissociation process. Based on the integrated P(E), we conclude that the yield of ground state O₂ exceeds 20% at 588nm. It is very interesting to note that the dominant products are formed with translational energy consistent with formation of $O_2(^1\Delta)$. In fact, if the data in Figs. 29 and 30 are examined carefully, we are very tempted to postulate that the observed structure results from a progression in O_2 ($^1\Delta$), with the most probable vibrational level being v=3, if most NO is produced in v=0. Alternatively the data might be interpreted to indicate that in fact the O₂ is entirely formed in its ground vibrational state, with the most probable vibrational level being 8. Based on the data alone, it is impossible to state with certainty which interpretation is correct. However, if the symmetries of the appropriate orbitals are considered, formation of ground state O₂ is strongly forbidden for dissociation constrained to C_{2v} . 9,28 Of course, we do clearly observe this channel, which will be favored by departure from C_{2v} symmetry in the transition state.^{9,28} However, it is expected that the potential energy barrier for formation of O_2 ($^1\Delta$) will be smaller than

for the ground state, based on correlation of the symmetry species involved.³ In light of this, and our data, it would be tempting to say that $O_2(^1\Delta)$ is dominant. We are currently considering this question in detail, based on the appropriate symmetry arguments.

The possibility that the NO is primarily in v=1 can be explored based on the observed data in Fig. 30. If O_2 is in the $^1\Delta$ state, then the predicted maximum translational energy for v' = 3-4 appears to differ considerably from the data. Thus, it appears that NO v' = 0 is likely to be dominant. Of course, we expect that NO will appear in a range of vibrational levels, particularly in view of the fact that the N-O bond length decreases somewhat from 1.24Å in NO₃ to 1.15 Å in the NO product. However, our data does indicate that it is mostly formed in low vibrational levels. Usually, the resolution of our experiments is not sufficient to provide information on the disposal of energy into rotational degrees of freedom of the products. However, in the present case, it is clear that rotational energy of the products must be small. Otherwise, the P(E) could not show the structure reported here. A small amount of rotational energy in the two product molecules is very reasonable in this process. Since NO₃ has D_{3h} symmetry, the impact parameter for the recoiling O_2 + NO products must be nearly zero in the dissociation process. Consequently, we expect to see very little rotational energy since there are no strong forces to provide the rotational excitation. The transition state requires considerable symmetric bending to bring the two oxygen atoms within close enough proximity to form an O=O bond. Even if the transition state is nonplanar, we do expect that nearly all of the energy will appear in O2 excitation. Due to the small

impact parameter in the half collision process, the NO will be formed with nearly zero rotational energy. Since total angular momentum must be conserved in the dissociation process, the O₂ molecule is "constrained" to also having very small angular momentum, but in the opposite direction. This explains why we are able to see structure in a process where so many product degrees of freedom would normally be expected to wash out the structure entirely.

There is a noticeable difference between our observations near the O-NO₂ dissociation threshold and the observations of Welge⁴¹ near threshold for formation of NO + O in photodissociation of NO₂. We find that the yield of NO₂ + O from ground state NO₃ increases rather gradually from 0 to 100% over the wavelength range 590-584nm, an energy range of ~175cm⁻¹. The quantum yield for NO + O in photodissociation of NO₂, on the other hand, apparently undergoes the same increase over an energy range of only 1-2 cm⁻¹. This behavior appears to be mostly attributable to the expectation that the NO₃ density of states near threshold will be far higher than in NO₂, leading to a much smaller rate for dissociation at energies near threshold.

The 594nm data is plotted in the same manner in Figs 31 and 32. Of course the 0.5 kcal/mole decrease in photon energy produces the corresponding shifts in the observed peaks. All of the arguments stated above also apply to the 594nm data. Perhaps the most important observation is that the relative contribution from ground state O₂ is clearly smaller at the lower photon energy, i.e. the fast component shows a smaller contribution. At first glance, it might seem strange that less ground state O₂ is formed with the lower energy photon. However, based on the calculations of Siegbahn, 9 and by

Boehm and Lohr,²⁸ we expect that the yield of ground state O₂ will be very sensitive to the degree to which C_{2v} symmetry is broken. At the lower photon energy, there is slightly less internal energy available. However, the probability that the molecule will achieve a configuration below the ground state potential energy barrier will be critically dependent on its vibrational excitation. It appears that this is the dominant reason why only 0.5 kcal/mole more energy leads to nearly a factor of 2 greater yield of ground state O₂. As expected, we see a larger yield of more internally excited O₂ at the longer wavelength. These considerations will be explored in detail in a forthcoming publication.⁴⁵

V. CONCLUSIONS

We have demonstrated that NO₃ undergoes some very interesting photodissociation dynamics upon electronic excitation in the visible. The observed wavelength dependence is easily explained in terms of kinetic competition between the possible dissociation mechanisms. At an excitation energy just below threshold for dissociation of ground state NO₃ to NO₂ + O, all the ground state molecules dissociate to NO + O₂. This situation is seen at 590nm. At this wavelength, those NO₃ molecules in the beam having any vibrational excitation ($\sim 50\%$) dissociate by the dynamically more favorable route to NO₂ + O. Although at least 10- 20% of the O₂ is in its ground state, (depending upon wavelength), there is good evidence that a substantial fraction is actually formed in the $^1\Delta$ state. Based on the observed wavelength dependence for the channels, and measurements of the NO₂ + O translational energy release, we conclude that D₆(O-NO₂) = 48.55 \pm 0.16 kcal/mole. By combining this with the known values

for $\Delta H_f(NO_2)$, and $\Delta H_f(O^3P)$, we calculate $\Delta H_f(NO_3) = 19.01 \pm 0.24$ kcal/mole (OK), and $\Delta H_f(NO_3) = 17.75 \pm 0.24$ kcal/mole (298K). Although this result does not agree with recent thermochemical measurements, it is in excellent agreement with a calculated value based on the most recent values for EA(NO₃) and $\Delta H_f(NO_3^-)$.

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Table 1: Calculated NO_3 Vibrational Populations Assuming D_{3h} Symmetry¹²

Mode	Vibrational Frequency ¹²	$N(\nu_i)/N(\nu=0)$	
	(cm ⁻¹)	(600K)	(300K)
$\overline{ u_1}$	1050	0.08	0.0066
ν_2	762	0.16	0.026
ν_3	1492 (2)	0.056	0.0016
ν ₄	363 (2)	0.84	0.35

Table II: Vibrational energies of NO and various electronic states of O₂.25

Molecule	დ₅(cm⁻¹)	ω _e x _e (cm ⁻¹)
NO (2II)	1904.04	14.1
$O_2(^3\Sigma_{g}^{-})$	1580.19	11.98
$O_2(^1\Delta_g)$	1483.5	12.9
$O_2(^1\Sigma^+_{g})$	1432.77	(11)

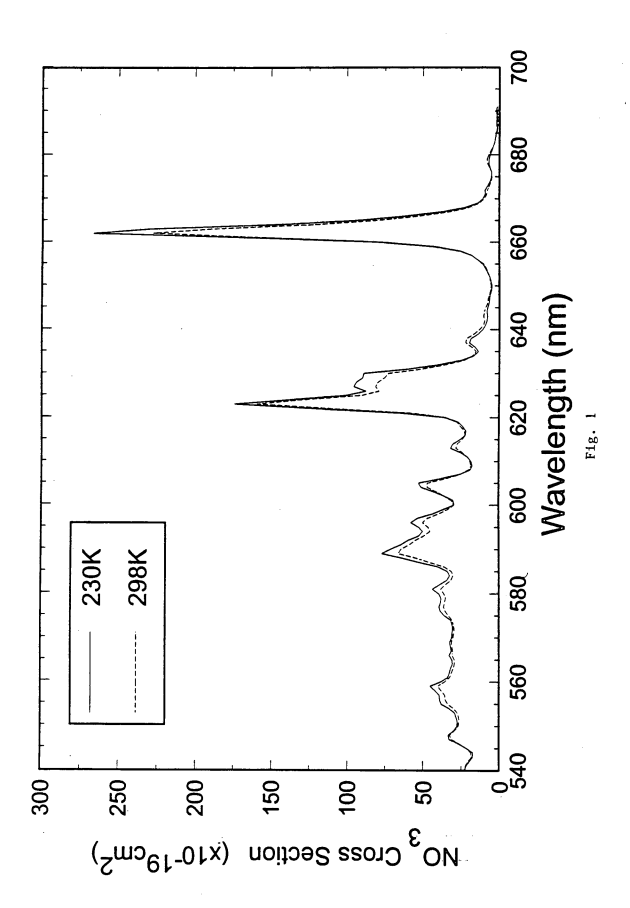
VIII. LIST OF FIGURES:

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- Fig. 2 Approximate Correlation and energy diagram for NO₃ photodissociation, obtained from Ref. 3.
- Fig. 3 NO and O quantum yields for NO₃ photolysis, as obtained by Magnotta and Johnston. Data reproduced from Ref 2b.
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 - B) Schematic of the improved nozzle design used on RSM
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- Fig.11 NO⁺ TOF spectra obtained at several detector angles from 590nm photolysis of NO₃.
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- Fig 32 Same as Fig. 31, but assuming NO(v=1).



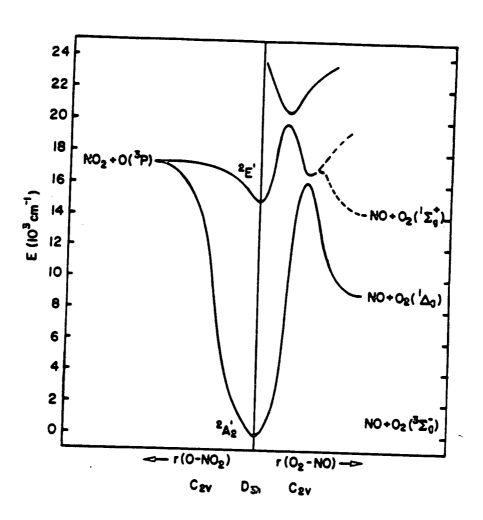
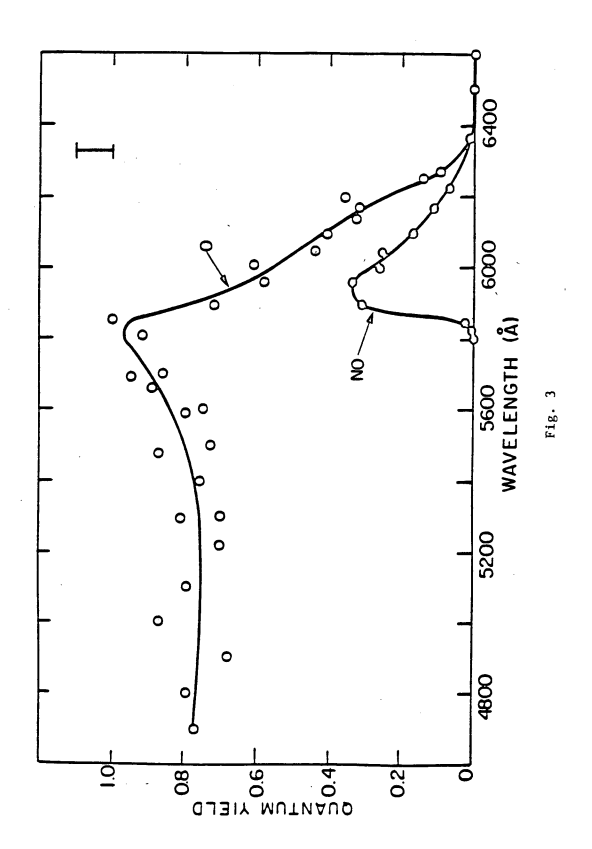


Fig. 2



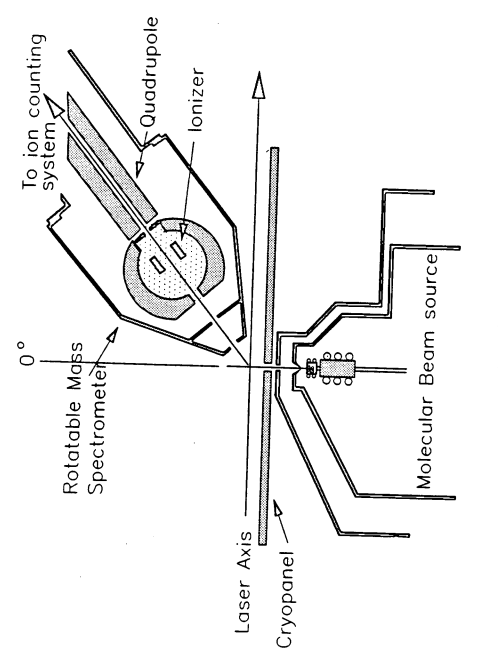
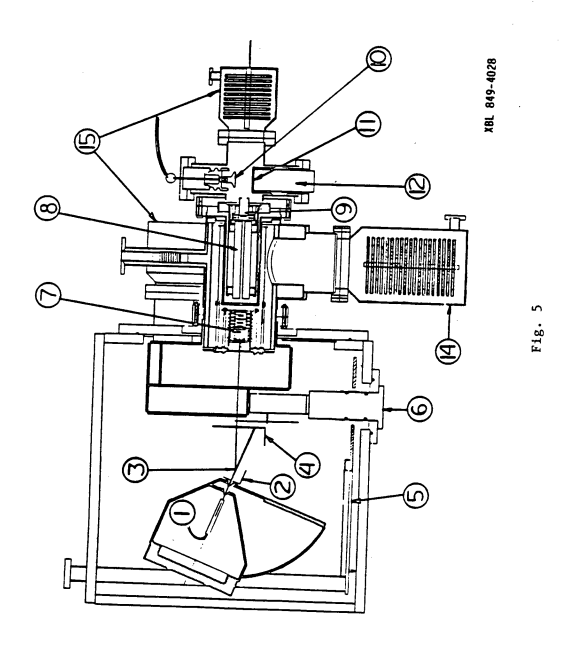
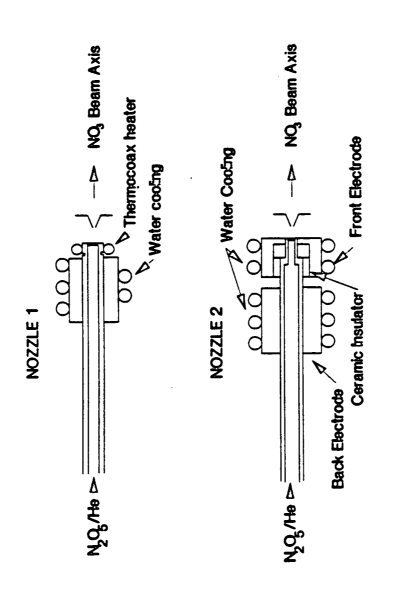
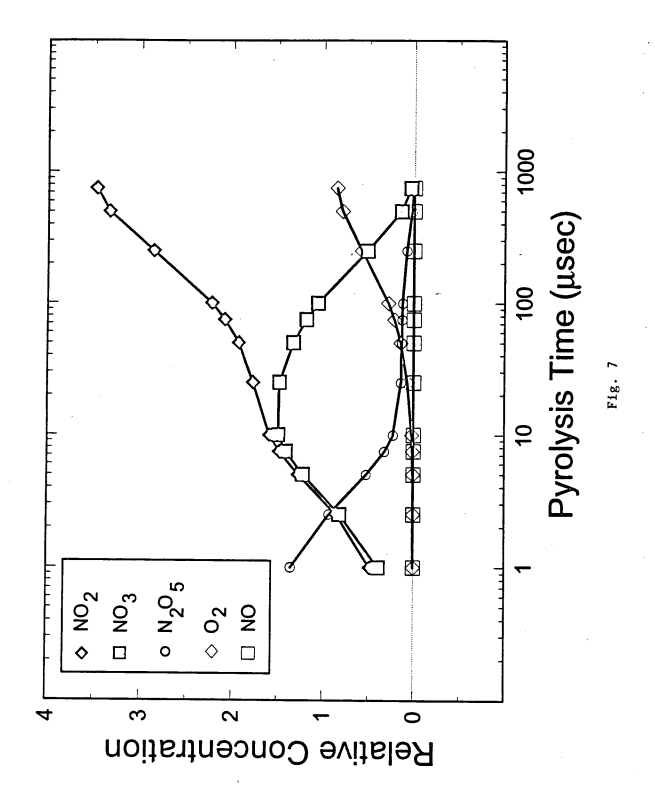


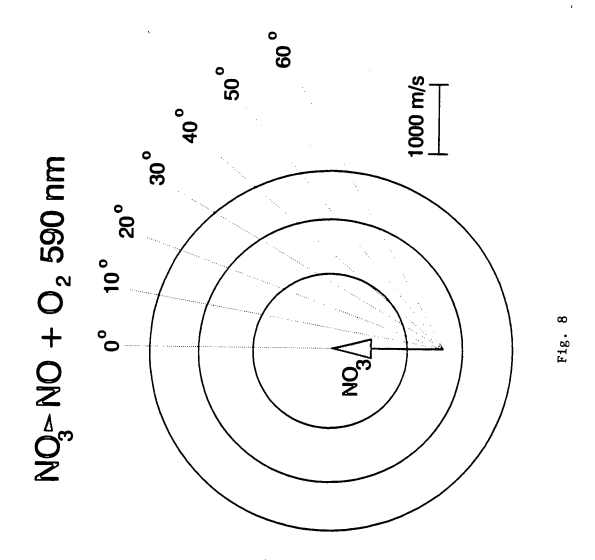
Fig. 4





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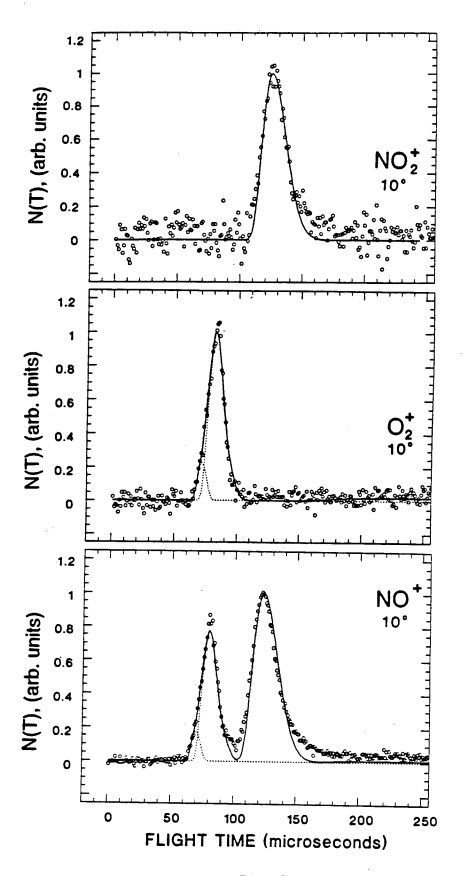
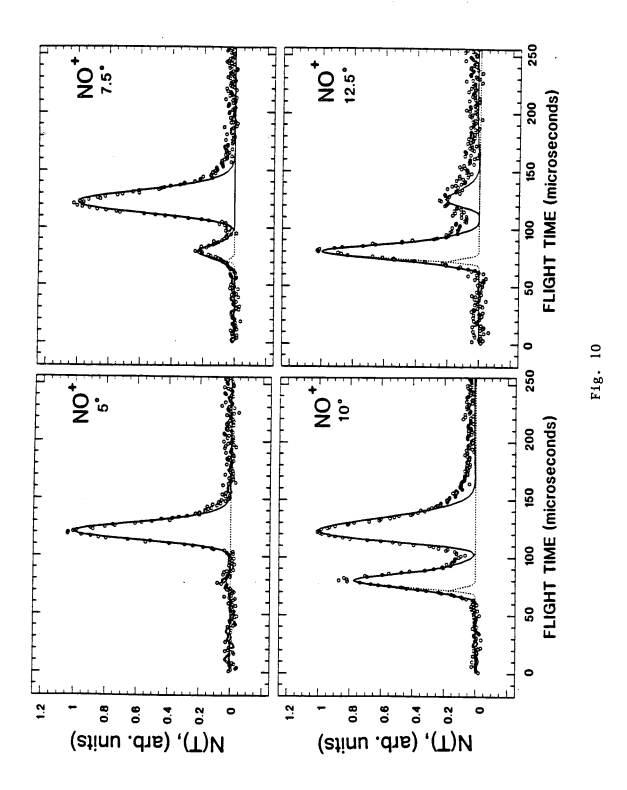
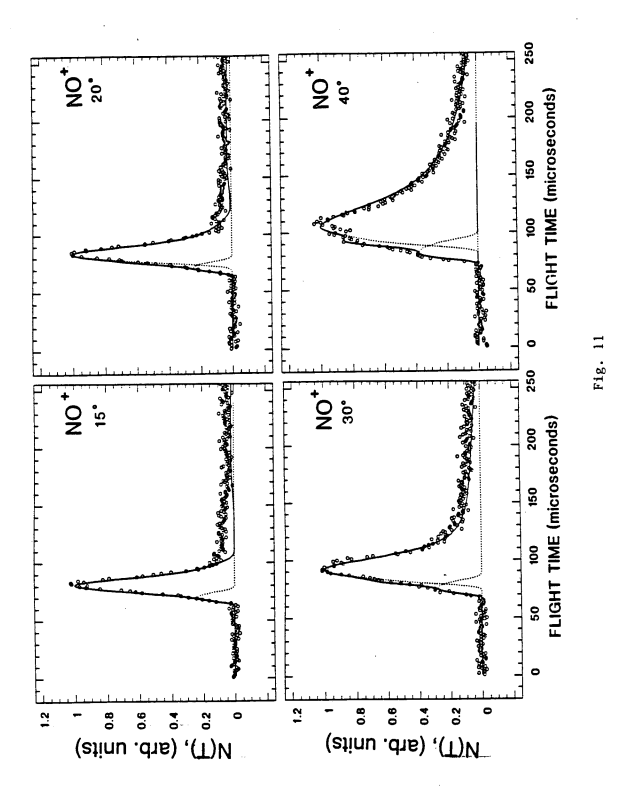
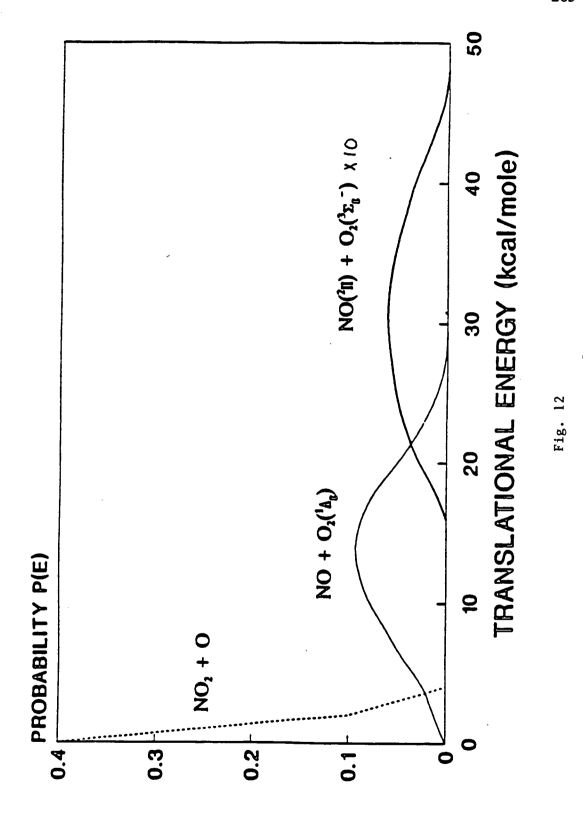
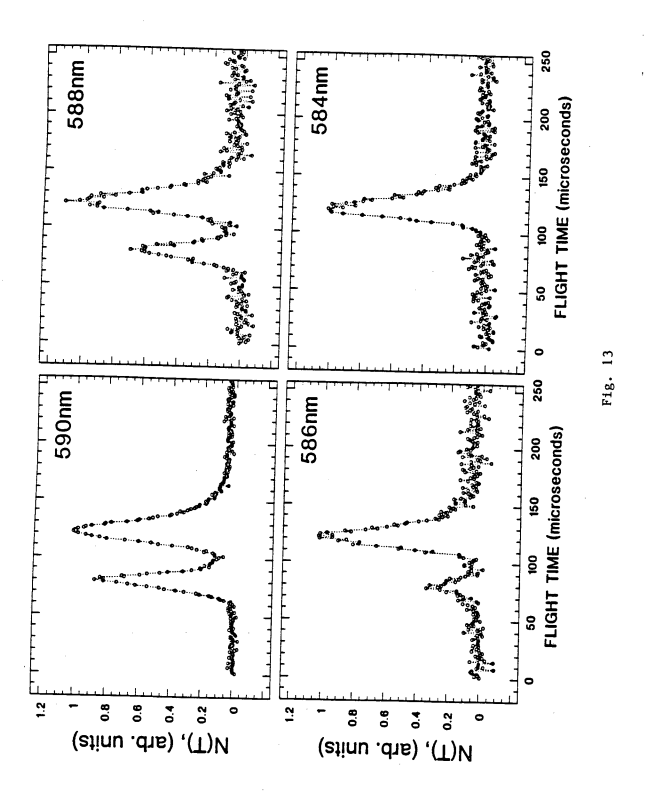


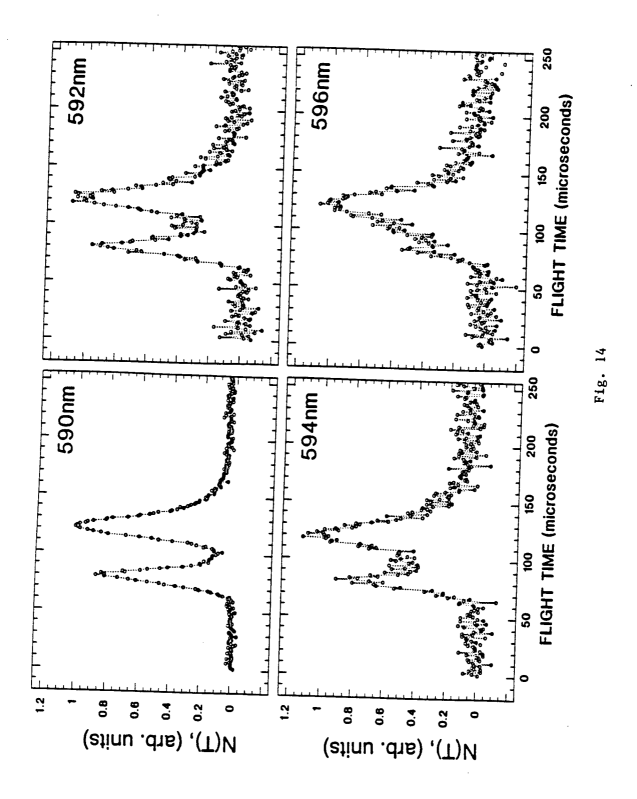
Fig. 9

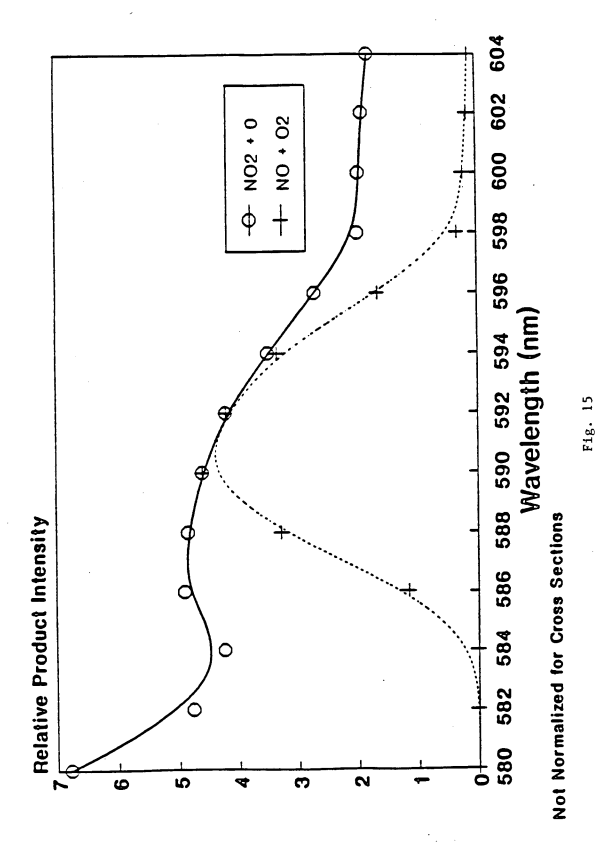


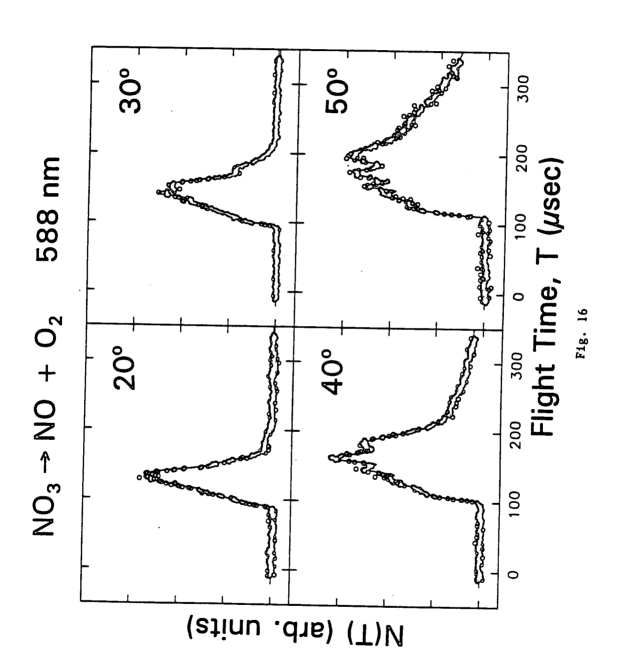


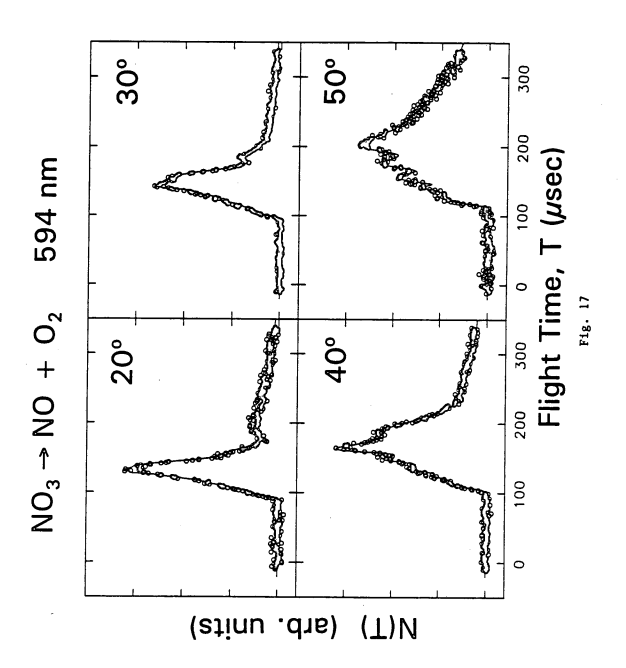


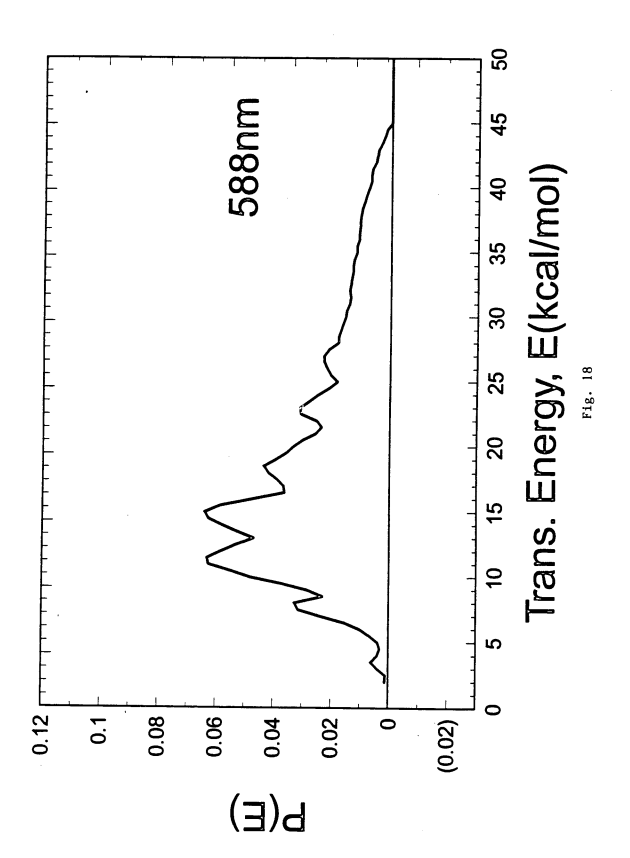


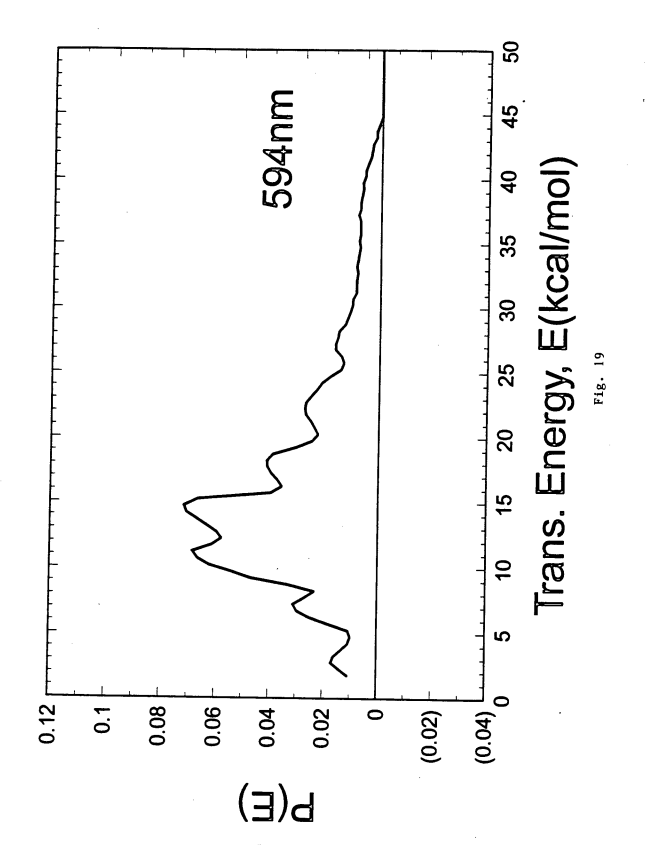












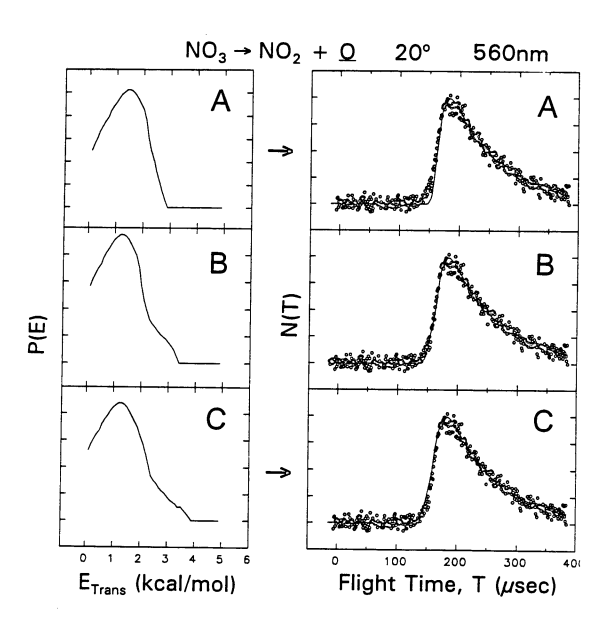


Fig. 20

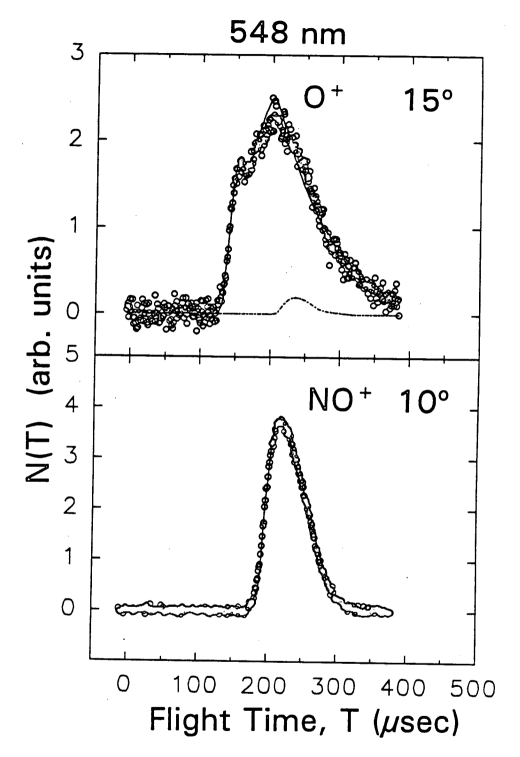
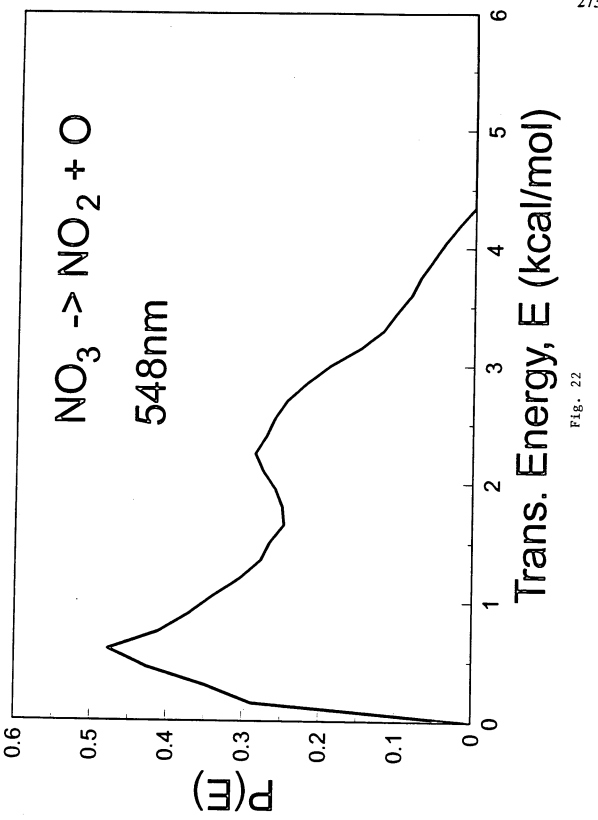
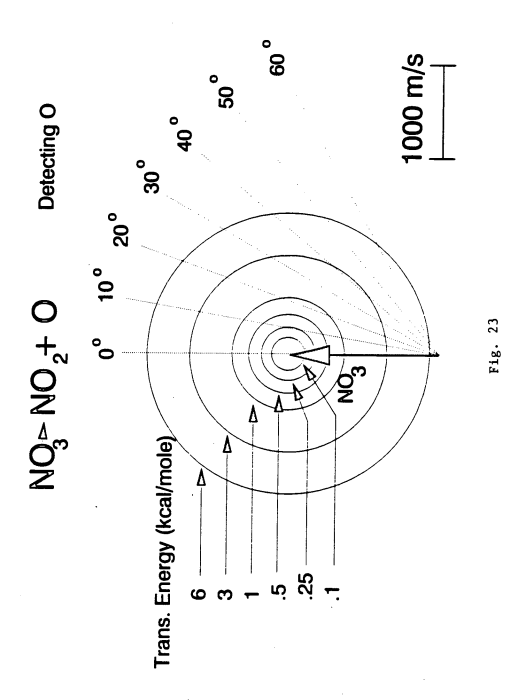
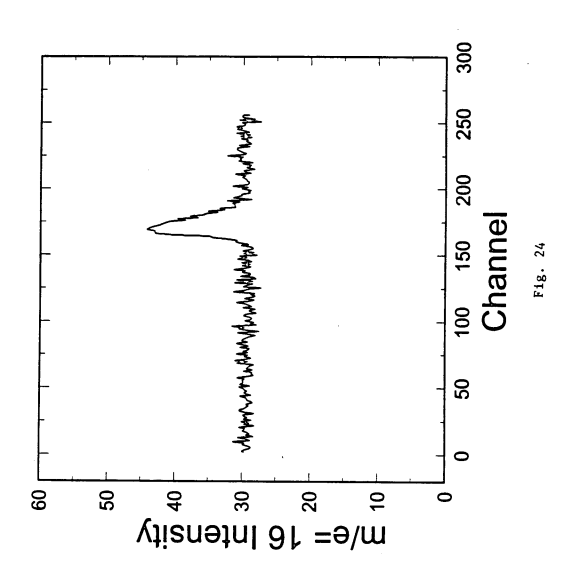


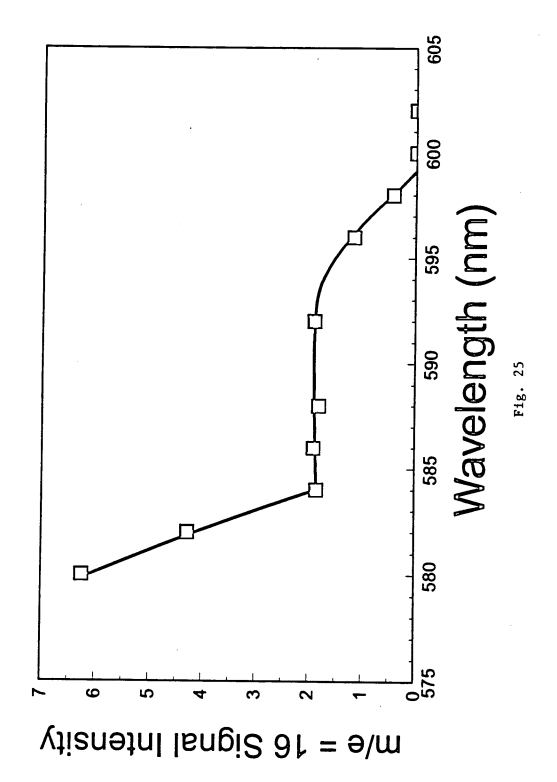
Fig. 21

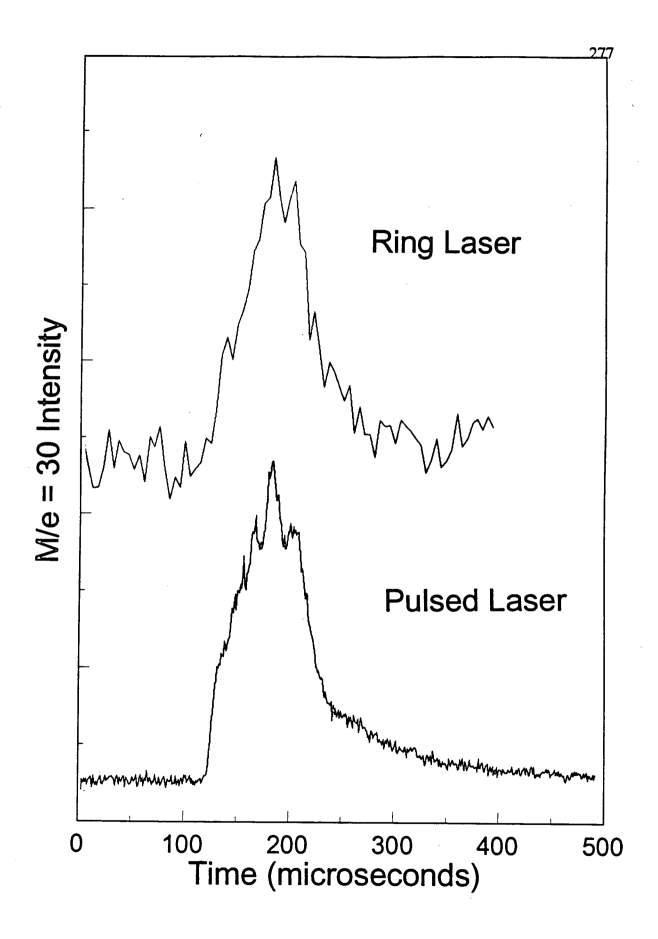


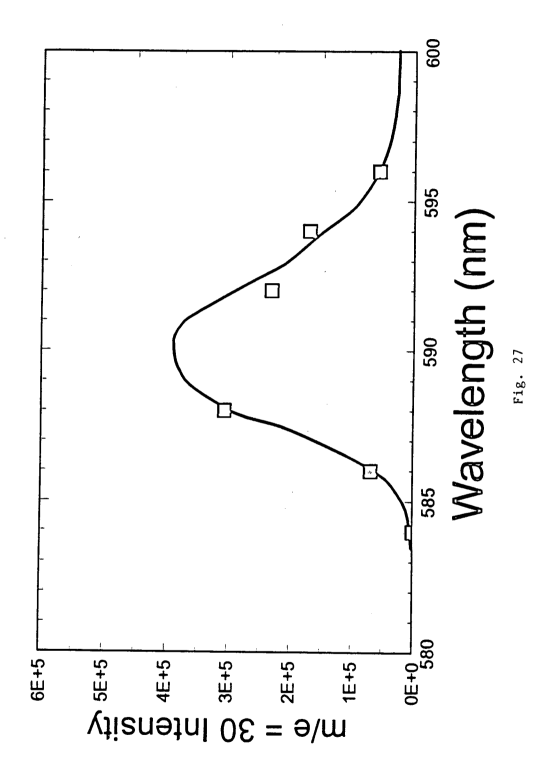


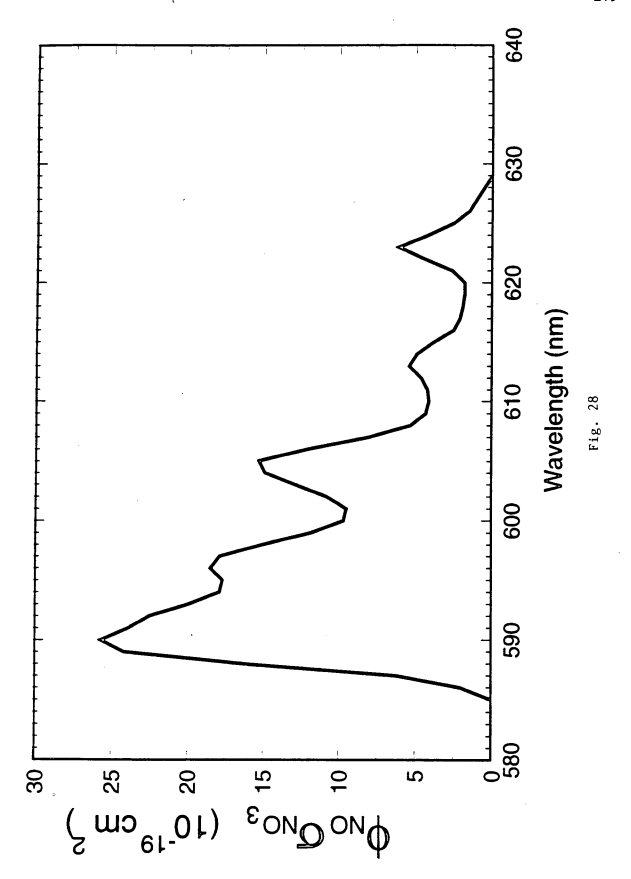


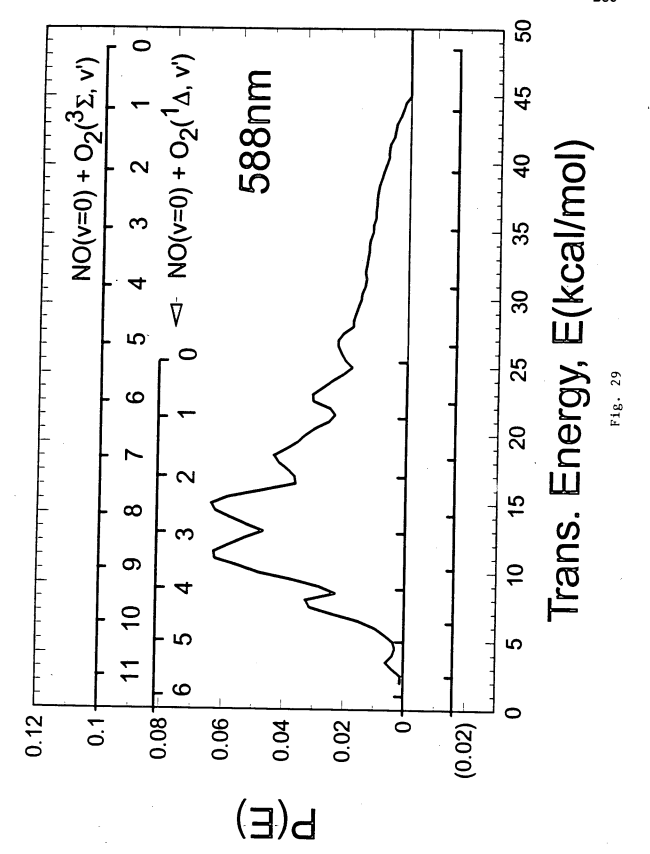


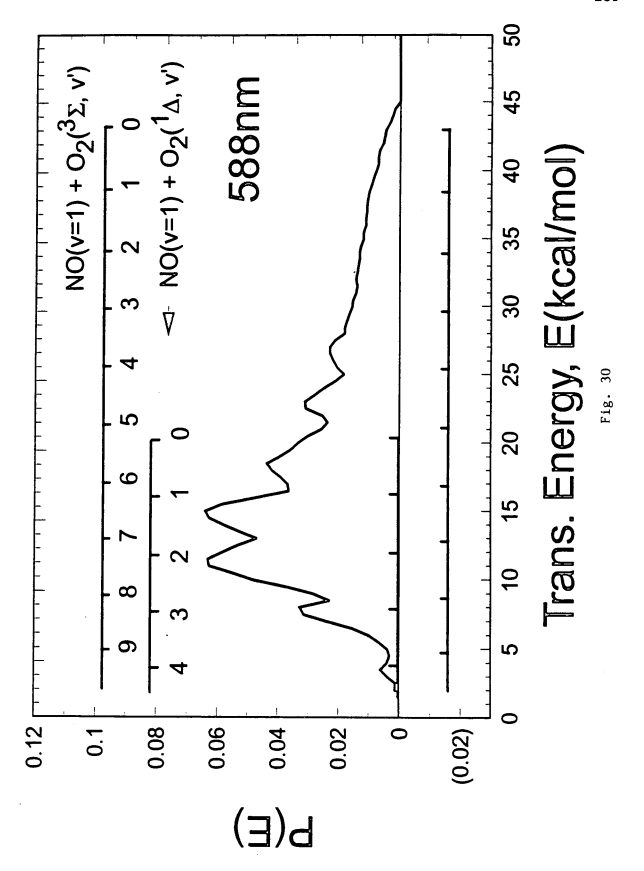


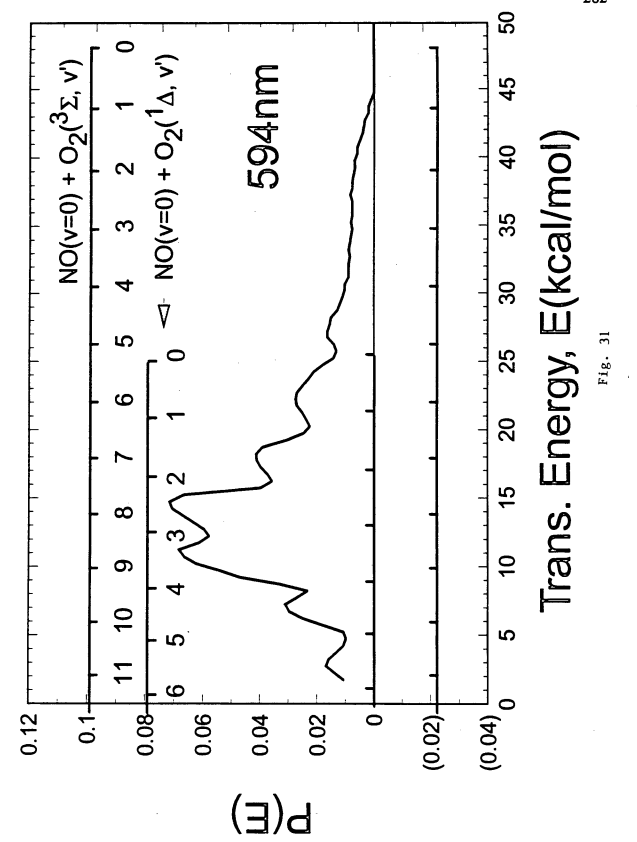


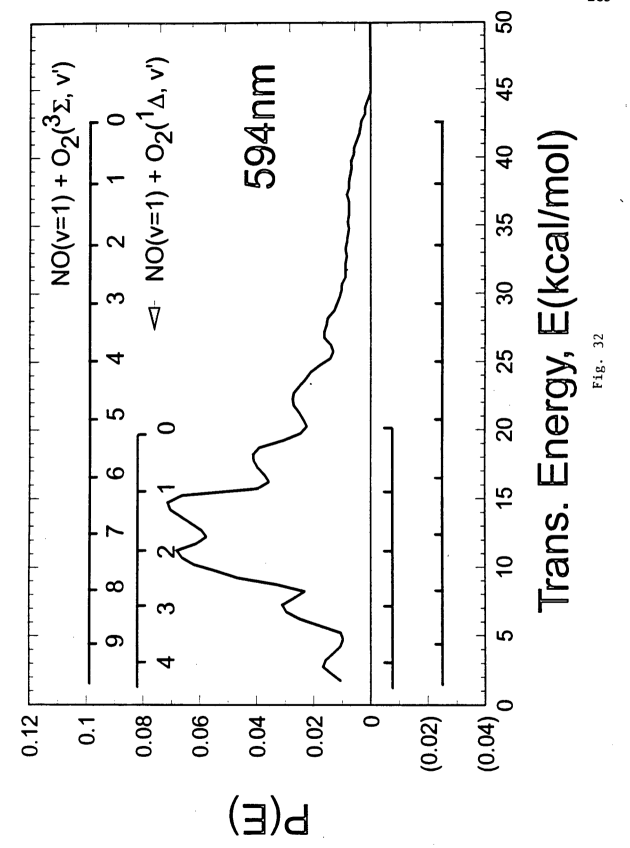












CHAPTER 5

Dynamics and Mode Specificity in OCIO Photodissociation

ABSTRACT

The photodissociation of OCIO in a molecular beam was studied using photofragment translational energy spectroscopy at wavelengths between 350 and 475nm. Although the dominant products are $ClO(^2\Pi) + O(^3P)$, we observe formation of $Cl(^2P)$ + $O_2(^3\Sigma_g)$, and find strong evidence for $Cl(^2P)$ + $O_2(^1\Delta_g)$. The total yield reaches a maximum of $3.9\pm0.8\%$ near 404nm and decreases to <0.2\% under the absorption peaks below 370nm. It is known that the OClO (A²A₂-X ²B₁) absorption spectrum has a well defined vibrational progression resulting from excitation to the $(\nu_1,0,0)$, $(\nu_1,1,0)$, $(\nu_1,0,2)$, and $(\nu_1,1,2)$ levels of the excited state. We find that the branching ratios for the two chemically distinct channels display mode specificity. Although the yield of Cl + O₂ from OClO with symmetric stretching $(\nu_1,0,0)$ excitation is substantial (3.2%), a modest enhancement (to 3.9%) is seen under neighboring peaks associated with symmetric stretching + symmetric bending $(\nu_1, 1, 0)$. However, the Cl + O_2 yield is suppressed by a factor of ~10 (to 0.4%) from states at nearly the same energy with asymmetric stretching excitation. The Cl + O₂ results from concerted unimolecular decomposition with release of up to ~ 68 kcal/mol in relative translational energy.

I. INTRODUCTION

Nearly two decades have passed since Molina and Rowland¹ first pointed out the possibility that reactions of photogenerated Cl atoms in the atmosphere could lead to catalytic destruction of global ozone. In 1985, Farman and coworkers² noted a considerable decline in the abundance of Antarctic ozone, with a decrease in total column ozone of up to 50% in the month of September. These observations have been substantiated in more recent years,³ and have led to a considerable increase in experimental activity; both in the field, as well as in the laboratory. It has been found that the concentration of ClO and OClO in the Antarctic stratosphere reach levels nearly two orders of magnitude higher than normal, and these levels strongly anticorrelate with O₃ abundance.⁴ Although a number of catalytic cycles have been implicated in ozone depletion,⁵⁻⁷ it appears that the two major mechanisms are:

$$2(Cl + O_3 \rightarrow ClO + O_2)$$

$$ClO + ClO + M \rightarrow (ClO)_2 + M$$

$$(ClO)_2 + hv \rightarrow ClOO + Cl$$

$$ClOO + M \rightarrow Cl + O_2 + M$$

Net: $2O_3 + hv \rightarrow 3O_2$ (Cycle 1)

and,

$$Cl + O_3 \rightarrow ClO + O_2$$

 $Br + O_3 \rightarrow BrO + O_2$
 $ClO + BrO \rightarrow Br + ClOO$
 $ClOO + M \rightarrow Cl + O_2 + M$

Net: $2O_3 + hv \rightarrow 3O_2$ (Cycle 2)

However, an alternative chemical product in reaction of ClO + BrO (cycle 2) is possible:⁸⁻¹⁰

$$ClO + BrO \rightarrow OClO + Br$$
 (3)

Although the branching ratio for channel 3 is known to be greater than 50%, 8.9 OCIO in the atmosphere had not been considered important in cycles involving catalytic ozone destruction because it is rapidly photolyzed in sunlight:

OCIO (X
$${}^{2}B_{1}$$
) \rightarrow OCIO (A ${}^{2}A_{2}$) \rightarrow CIO(${}^{2}\Pi$) + O(${}^{3}P$). (4)

There have been no experimental measurements of the quantum yield for reaction (4) in the gas phase. However, it has been taken as unity in atmospheric modelling calculations.⁶ If this quantum yield is correct, the formation of OCIO and its subsequent photolysis regenerates free O atoms. Since they rapidly recombine with O₂ to regenerate

O₃, the OClO cycle would have no effect on ozone concentration in the atmosphere.

In 1990, Vaida and coworkers¹¹ reported experimental evidence for the following reaction under collision free conditions:

OCIO (X
$${}^{2}B_{1}$$
) \rightarrow OCIO (A ${}^{2}A_{2}$) \rightarrow OCIO(${}^{2}B_{2}$) \rightarrow CIOO \rightarrow CI + O₂* (5)

In their experiment, a molecular beam containing OClO was crossed with the output of a focussed dye laser operating at wavelengths near 362nm. The OCIO was photolyzed and the products were subsequently ionized by nonresonant absorption of additional photons. These ions were extracted into a time-of-flight mass spectrometer, and their intensities monitored as a function of laser wavelength. They observed a feature in the Cl atom REMPI spectrum at 362nm that they attributed to isomerization of OCIO to ClOO, followed by dissociation to Cl + O₂. In a subsequent report to Nature, ¹² they estimated that if the quantum yield for Cl + O₂ is 100% over its entire absorption spectrum, the gas phase photoisomerization mechanism could lead to up to 30% of the O₃ depletion seen in Antarctica. If the quantum yield was only 10%, then it would still have an appreciable effect on the overall budget of O₃ in the Antarctic atmosphere, and should be included in atmospheric modelling. The notion of a gas phase isomerization mechanism was largely inspired by an early observation of ClOO in matrix isolation experiments by Arkell and Schwager. 13 They found that ClOO was readily formed in the matrix by photolysis of OClO at wavelengths near 365nm. Apparently, photolysis at other wavelengths led to a smaller yield of ClOO. Matrix assisted photoisomerization of

OCIO seemed to provide a much more efficient source of ClOO than other methods, such as photolysis of $Cl_2 + O_2$ mixtures. Gole has performed calculations on the ground and electronically excited states of OCIO and ClOO.¹⁴ He postulated that coupling between these electronically excited states of the two species provides a likely mechanism for photoassisted rearrangement of OCIO to ClOO in the matrix.

The ideas derived from the matrix isolation work have been extended to the gas phase in recent years, 11,12,16,18 with some controversy regarding the quantum yields for Cl + O_2 . 11,15,16 The quantum yield of Cl atoms was not determined in the original work of Vaida, and coworkers. 11 However, Lawrence, *et. al.*, 15 reported that the quantum yield of Cl was below 5 x 10^4 between 359 and 368nm, and that the Cl + O_2 channel had a negligible impact on the balance of atmospheric ozone. Quite recently, Bishenden and coworkers 16 disagreed, concluding that Cl atoms are formed upon single photon excitation of OClO near 362nm with a quantum yield of 0.15 ± 0.1 . The Cl atom yield at other wavelengths was not explicitly reported, but a structured Cl REMPI action spectrum was presented 16 for the wavelength range 355nm-367nm. Although it is now apparent that heterogeneous processes contribute to the depletion of ozone, 5 a clear understanding of the photochemistry of atmospheric molecules under collision free conditions is required before addressing other complicating factors, such as perturbations by solvent molecules or adsorption on ice crystals.

A number of factors inspired the present experiment. First, the NO_3 experimental work (Chapter 4) had already been completed and the analysis was in progress. The observation of a significant yield of $NO + O_2$ in that system prompted a search in the

literature for other previously studied molecules exhibiting similar behavior. It was hoped that comparisons with NO₃ might lead to additional insight into this highly complex molecule. Indeed, OCIO appeared to be such a system, ¹¹ but the experimental evidence for the formation of Cl + O₂ in the gas phase was not compelling, as will be discussed below. The second motivation was that the reaction cross section for Ba + OCIO \rightarrow BaCl + O₂ was found to appreciable (Chapter 3). This reaction involves concerted central atom abstraction, and is initiated by long range electron transfer from Ba to the OCIO 2b₁ orbital, leading to a decrease in OCIO bond angle and increase in ClO bond length. ¹⁴ The situation is quite analogous to OCIO (A²A₂ \leftarrow X ²B₁) excitation, since it also involves promotion of an electron to the 2b₁ orbital, leading to similar changes in molecular geometry favorable to formation of O₂. ¹⁴

The photodissociation dynamics of the bent triatomic molecules O_3 and NO_2 has been studied extensively over the past 20 years using a wide range of techniques. Triatomics are particularly amenable to detailed study because the parent molecule possess only three vibrational degrees of freedom and their atom + diatom products are easily detected experimentally, and modelled theoretically. However, the apparent simplicity of O_3 and NO_2 is belied by a complex electronic structure and fast predissociation processes, leading to poorly understood features in their electronic absorption spectra. The recent work of Vaida and coworkers¹⁷⁻¹⁹, on the absorption spectroscopy of OClO has led to a reinterpretation and considerable clarification of the OClO $(A^2A_2\leftarrow X^2B_1)$ absorption spectrum. Their work led to accurate assignments of the line positions, making it possible to use a tunable dye laser to prepare the OClO molecule

in well defined vibrational levels of the electronically excited (A²A₂) state.

Figure 1 shows the energetics of OCIO decomposition.²⁰ At all wavelengths, three electronic states of O₂ are thermodynamically possible. In none of the experiments reported to date^{11,16} was the electronic state of the O₂ determined. In fact, in no case was O₂ directly observed—all experimental evidence has been based on observation of the Cl atom alone. This leaves open the possible involvement of other sources of Cl signal, such as photodissociation of the dominant ClO product.

Using the technique of photofragment translational energy spectroscopy,²³ we find that $Cl + O_2$ is formed from single photon absorption by OCIO, but the yield is only appreciable (>1%) at wavelengths *longer* than 390nm. Unlike the previous measurements,^{11,16} we have observed both momentum matched $Cl + O_2$ products, and the quantum yields are based on direct measurements under well defined single photon conditions. Translational energy measurements show that O_2 is definitely formed in the ground $\binom{3}{2}$ state and there is very strong evidence for a substantial fraction in the first excited $\binom{1}{4}$ state.

Spectroscopic work by a number of groups^{17,18,21} has clearly shown that the OCIO absorption linewidths increase upon tuning the excitation source to the blue. This is a clear signature of a decreasing excited state lifetime, possibly by faster predissociation via the "dark" ²B₂ state. ^{17,18,21} Since the ²B₂ state cannot be accessed by a single photon transition from the ground state, it has not been observed experimentally. ¹⁷ However, ab initio calculations¹⁴ indicate that it probably lies at energies lower that the ²A₂ state, which is the only excited state to be observed experimentally. The linewidths associated

with excitation of asymmetric stretching + symmetric stretching (ν_1 ,1,0), or symmetric bending + symmetric stretching (ν_1 ,0,2), are known to be considerably broader than those due to symmetric stretching excitation (ν_1 ,0,0) alone.^{17,18,21}

The OCIO molecule has a structured absorption spectrum^{17,22} corresponding to progressions in $(v_1,0,0)$, $(v_1,1,0)$, $(v_1,0,2)$, and $(v_1,1,2)$ of the excited 2A_2 state (Fig. 2). Despite a dissociation lifetime of many vibrational periods, 17,18,21 we do observe mode specificity in the branching ratios for the two channels. The yield for Cl + O_2 is always greatest (up to 3.9%) under OCIO absorption peaks associated with combinations of symmetric stretching with symmetric bending $(v_1,1,0)$ motion. Although the yield is only slightly smaller from pure symmetric stretching modes $(v_1,0,0)$, excitation at nearly the same energy to a symmetric stretching + asymmetric stretching mode $(v_1,0,2)$ leads to a tenfold decrease in the Cl + O_2 yield. Mode specific behavior is also seen in the ClO vibrational energy distributions which are strongly dependent on the initial vibrational state of the OCIO molecule.

II. EXPERIMENTAL

The experiments were conducted using a molecular beam apparatus equipped with a rotatable molecular beam source and fixed detector.²³ The unfocussed output from an excimer-pumped Lambda-Physik FL2002 dye laser was crossed at 90 degrees with a seeded supersonic molecular beam containing 10-20% OCIO in helium or argon. In some experiments, a continuous beam was formed by expanding ~150 Torr of the mixture through a 0.007" dia. nozzle heated to ~80°C. Experiments were also conducted using

a room temperature pulsed nozzle (0.020" dia.) and a pressure of 1 atm. The OClO was generated *in-situ* by flowing a 5-10% Cl_2 /inert gas mixture through a U-tube packed with glass beads and NaClO₂, as described elsewhere.²² The Cl_2 impurity in the beam, measured using a mass spectrometer, was found to be negligible provided that the NaClO₂ is not too dry-- apparently the reaction $Cl_2(g) + 2NaClO_2(s) \rightarrow 2NaCl(s) + 2ClO_2(g)$ requires the presence of some moisture.

A small fraction of the photodissociation products recoiled away from the molecular beam axis into an electron bombardment ionizer located 36.7 cm away from the interaction region.²³ Ions were extracted into a quadrupole mass filter and detected by a Daly ion detector. A multichannel scaler, triggered by the laser pulse, was used to record the product time of flight spectra at various angles between the molecular beam and detector. The product time of flight spectra were analyzed using an updated version of the forward convolution program CMLAB2, as described previously.²³

III. RESULTS AND DISCUSSION

Under all absorption peaks between 350nm and 475nm, the dominant products (>96%) from dissociation of OCIO are $ClO(^2\Pi) + O(^3P)$. Product time of flight (TOF) spectra were recorded at Cl^+ , ClO^+ , O^+ , and O_2^+ . We observed Cl^+ daughter ions from fragmentation of ClO (from ClO + O) in the electron bombardment ionizer, and Cl^+ parent ions from the $Cl + O_2$ channel. As predicted from the Newton diagram (Fig. 3), both photodissociation channels can be seen by monitoring Cl^+ and can be distinguished by their TOF spectra. Even if the O_2 fragment is formed in v=5 of the $^1\Sigma^+$, state (the

second electronically excited state), the recoil velocity of its Cl atom partner will be greater than the velocity of ClO (v=0) from the ClO + O channel.

Time of flight spectra recorded at Cl⁺ (m/e=35) are shown in Figure 4 for three neighboring excitation wavelengths, corresponding to the (5,1,0), (4,0,2), and (5,0,0)vibrational levels of OClO (A²A₂). The angle between the molecular beam and detector was 20°, and a 20% OCIO/He mixture was expanded through a room temperature pulsed nozzle. The slower peak in the TOF spectrum was identical to that seen in the ClO+ TOF and results from ClO fragmentation in the ionizer. The fast peak is due to Cl from the Cl + O₂ channel. Our assignments are confirmed by observation of the respective momentum matched O and O₂ fragments (Fig. 5). Based on linewidth measurements^{17,18}, it is known that the OClO dissociation lifetime is many picoseconds at these wavelengths. Thus, unless the laser was tuned very close to the origin¹⁷ of a vibrational band, we found that rotating the laser polarization using a double fresnel rhomb had no effect on product intensities, since the parent molecule rotated at least several times before dissociating. All data reported here was obtained with the dye laser tuned 4-5 cm⁻¹ to the red of the bandhead positions reported in Ref. 17. Although our laser does not provide rotational resolution, ($\Delta E \sim 0.2 \text{cm}^{-1}$), we see only minor effects on the shapes of the TOF or Cl + O₂ yields upon tuning across the rotational envelopes of given vibrational levels of OClO.

The translational energy distributions for both channels from excitation of the (5,1,0) band are shown in Figure 6. They are based on data shown in Figures 4 and 5, as well as other data (not shown) obtained at other angles between the molecular beam

and detector.²⁴ The fastest component of the Cl + O₂ channel, with E_{trans} up to ~68 kcal/mole, must correspond to formation of ground state O₂ ($^3\Sigma_g$). The additional feature peaking at ~33 kcal/mole is very strongly suggestive of formation of excited O₂ ($^1\Delta_g$). As shown in Fig. 4, the shapes of the Cl + O₂ times of flight (and O₂ internal state distributions) are not strongly dependent on the initial OClO vibrational level. The ClO internal state distribution, on the other hand, is very sensitive to the initially prepared OClO state. Structure due to vibrational excitation of the ClO($^2\Pi$) product is quite well resolved, particularly when its O atom recoil partner was monitored (Fig. 5). Analysis is currently under way to extract approximate vibrational energy distributions as a function of OClO vibrational state.²⁴

The branching ratios for formation of Cl+O₂ relative to ClO+O were calculated from the fits to the experimental data at m/e=35(Cl⁺), accounting for the different Jacobian factors for the LAB+CM transformation for each channel. One additional factor that is required in the analysis is the relative detection sensitivity for ClO and Cl at m/e=35 (Cl⁺). This quantity was determined experimentally in a separate experiment. We monitored the equal yields of Cl and ClO from ClOCl photodissociation at 423nm. ^{24,25} The fast peak in the m/e=35 TOF resulted from Cl and the slower peak was from fragmentation of ClO-- their contributions led to a very accurate measure of the relative detection sensitivity for the two fragments. ²⁶ The measured branching ratios were converted to absolute yields by assuming that $\Phi_{diss} = \phi_{Cl+O2} + \phi_{ClO+O} = 1.0.$ Our absolute Cl + O₂ yields should be accurate to $\pm 20\%$.

Our results indicate that the yield for $Cl+O_2$ has a maximum of 3.9 \pm 0.8% under the peak near 404nm dropping to <0.2% under peaks at wavelengths below 370nm. The $Cl+O_2$ channel is observed at longer wavelengths, but is weaker—the yield under the (4,0,0) band at ~423nm is ~1.2%. Our conclusion that the yield is negligible below 370nm is in qualitative agreement with Lawrence, ¹⁵ who could not actually see $Cl+O_2$, but placed an upper limit of 0.05% below 368nm. Extensive averaging would be required in our experiment to further reduce our upper limit of 0.2% for λ <370nm. All previous experimental efforts ^{11,15,16} have focussed on a wavelength range near 365nm, apparently because early matrix isolation experiments ¹³ found that the formation of ClOO by irradiation of OClO was favored using filters transmitting at that wavelength. We have not been able to observe any $Cl+O_2$ in this region in our molecular beam experiment.

Bishenden, et al, ¹⁶ reported a REMPI spectrum with the peak Cl intensity near 362nm. This appeared surprising since the OCIO molecule has an absorption *minimum* at this wavelength. However, these authors have recently explained this behavior in terms of an accidental laser calibration problem. The spectrum should be shifted by 2nm to the blue, so that it resembles the OCIO absorption spectrum. However, their shifted data would still indicate a Cl + O₂ quantum yield in excess of 10%, which is at least 50X higher than our upper limit of 0.2% in this wavelength region. It seems quite possible that the observed Cl signal results from a multiphoton process involving dissociation of OCI by the REMPI laser. This would not be surprising since the focussed laser at 235nm will be absorbed strongly by the nascent ClO molecule, and their laser

power dependences could be misleading due to saturation. The Cl atom product is then ionized by the 2+1 REMPI process.

It is important to note that our measurement of the $Cl + O_2$ quantum yields do not require knowledge of the OCIO absorption cross section. This is one significant advantage to the present technique-- virtually all other methods require that this quantity be known. In the case of OCIO, there is considerable rotational structure under its vibrational envelopes. This structure will be resolved to some extent if dye lasers having linewidths less than $1 cm^{-1}$ are used, and this will lead to considerable uncertainty in the measurement. Our calibration is only based on the relative sensitivity of our detector to Cl and ClO. This was accurately determined by measuring the signal levels from equal numbers of these products from photodissociation of ClOCl.

It is interesting to compare our gas phase results with the matrix results of Arkell, et al. We observe a significant yield (>1%) of Cl + O₂ only at wavelengths longer than 390nm, whereas they found that ClOO is favored in the matrix by irradiation at \sim 365nm. This *could* suggest that our observation of the Cl + O₂ channel under isolated gas phase conditions might be unrelated to observation of ClOO in the matrix experiments. Photodissociation near 365nm could lead to formation of ClO + O in the matrix, but due to the matrix cage effect, back reaction of the nascent O atom leads to ClOO. Back reaction in the matrix cage will be facile because the ClO(2 II) + O(3 P) electronic states correlate directly with ClOO (A',A"). The "enhancement" in yield of ClOO in the matrix by irradiation near 365nm, (rather than at 220nm < λ < 350nm or 450nm < λ < 590nm) could simply result from the OClO absorbing most strongly in

It is also possible that the 3000cm⁻¹ shift between the wavelength maxima for the gas phase Cl + O₂ channel and the matrix ClOO channels is real. Although 3000cm⁻¹ is a very large shift, it may not be unreasonable since the OClO (A²A₂-X ²B₁) transition does have considerable charge transfer character, and the crossings of the relevant potential energy surfaces could be shifted significantly in the presence of a matrix host. While it is likely that a fraction of the ClOO in the matrix does result from recombination of ClO + O due to the cage effect, Gole has noted a number of interesting features in the matrix work which have been taken to result from "matrix assisted photoisomerization." Our results indicate that comparisons between gas phase and condensed phase behavior will be complex in the case of OClO photochemistry.

Previous discussion regarding OCIO \rightarrow Cl + O₂ in the gas phase has centered on a photoisomerization mechanism involving an intermediate ClOO radical. ^{11,12,16-18} However, a mechanism formally involving isomerization of OCIO to ClOO in the gas phase, followed by simple bond rupture does not appear to be consistent with our observations. The large translational energy release that we observe in the recoiling Cl + O₂ is characteristic of a concerted unimolecular decomposition from a highly constrained transition state, followed by strong repulsion between the products. This transition state is probably best considered to be an OCIO molecule with a strongly compressed bond angle and only some distortion from C_{2v} symmetry. This is consistent with the Cl-O antibonding and O-O bonding nature of the 2b₁ orbital into which the electron is excited, resulting in a decrease in OCIO bond angle from 117 to ~ 107°, and

increase in Cl-O bond length from 1.47 to 1.62Å. Our observation that symmetric stretching + symmetric bending excitation promotes the Cl + O₂ channel strongly supports a mechanism involving concerted decomposition from a transition state geometry close to C_{2v} . The region of the Cl + O₂ potential energy surface corresponding to the Cl-OO isomer represents a very shallow minimum with $D_o(\text{Cl-OO}) = 4.76 \pm 0.49 \text{ kcal/mol}^{32}$ and $\Theta_{\text{Cl-OO}} \sim 110^{\circ}.^{13.14}$ In the absence of a large torque on the O=O molecule, the newly born Cl atom will be near the C_{2v} axis of OClO, perpendicular to the O=O bond axis. Since the Cl-O₂ interaction at this angle is repulsive, ¹⁴ the Cl-O₂ "isomer" will not exist for much more than one vibrational period. Instead, strong product repulsion leads to a large Cl +O₂ recoil energy of up to $\sim 68 \text{ kcal/mol}.$

The dissociation to CIO + O is thought to occur by coupling between the prepared ${}^{2}A_{2}$ and "dark" ${}^{2}B_{2}$ state. The observation that the OCIO absorption linewidths are independent of rotational energy has been interpreted in terms of a mechanism involving spin-orbit coupling between these states. The coupling is expected to be enhanced considerably by asymmetric stretching and bending motion of the OCIO molecule. The importance of asymmetric stretching in promoting the CIO + O channel is certainly not surprising in view of the fact that it is the reaction coordinate for the process. Indeed, for a given excitation energy, the OCIO absorption lines are known to be a factor of \sim 7-8 times broader (i.e. \sim 7-8 times faster dissociation) when excitation is accompanied by 2 quanta of asymmetric stretching. Interestingly, excitation of asymmetric stretching decreases *both* the excited state lifetime, *and* the CI + O_{2} -yield by roughly the same factor. Based on the known linewidths, The OCIO

dissociation lifetime is extraordinarily long for a triatomic molecule (>10ps), even at these energies which are >10kcal/mol above the ClO + O threshold. This is indicative of very weak coupling between the initially prepared 2A_2 state and that responsible for production of O + ClO. Although electronic excitation leads to strong symmetric motion, a sparse excited state level density with widely different frequencies for the three vibrational modes leads to very inefficient redistribution of energy into the reaction coordinate, asymmetric stretching. Consequently, only two quanta of asymmetric OClO vibration leads to a tenfold decrease in excited state lifetime. Our observation that the Cl + O₂ channel decreases with asymmetric stretching by a factor of 10 is most easily explained by faster decomposition to ClO + O, at the expense of the Cl + O₂ channel.

Based on the results presented in Fig. 7, and the measured OCIO absorption linewidths, we can make several conclusions regarding the role of symmetric bending in promoting the O + CIO channel. At all wavelengths where we observe $Cl + O_2$, $(380 \text{nm} < \lambda < 430 \text{nm})$, the quantum yield is enhanced only slightly when excitation is accompanied by symmetric bending, whereas the yield decreases dramatically ($\sim 10 \text{X}$) when accompanied by asymmetric stretching (Fig. 4). Since electronic excitation already results in a large decrease in OCIO bond angle, one additional quantum of symmetric bending apparently has a relatively minor additional effect on the $Cl + O_2$ quantum yield. However, it is known that the OCIO absorption linewidths are $\sim 4-5$ times broader when symmetric stretching is accompanied by one quantum of symmetric bending. Taking this observation with our finding that symmetric bending only leads to a modest enhancement of the $Cl + O_2$ yield, we conclude that symmetric bending leads to a substantially

increased rate of predissociation to OCl + O. This appears to be consistent with previous assertions 14,17,18,21 that it is the dark 2B_2 state that is involved in the OCl + O channel. The prepared 2A_2 and the dark 2B_2 states represent a Renner-Teller pair, becoming orbitally degenerate $^2\Pi$ states in the linear configuration. 17,18 Consequently, the coupling between these states will be very sensitive to the bending coordinate. This appears to be born out by our experimental results. Analysis of OCl vibrational energy distributions from a range of initially prepared OClO vibrational states is currently under way. We hope that this data will provide detailed insight into the nature of the interaction between the 2A_2 and 2B_2 electronic states.

As already stated, our observations strongly suggest that the transition state for the Cl + O_2 channel is likely to be near C_{2v} . The region of the Cl + O_2 interaction potential corresponding to the weakly bound Cl-OO isomer will not likely be accessed to a great extent in the gas phase dynamics. Of course, the dissociation process does involve coupling between the excited states of OClO and the A' and A" states of the Cl + O_2 interaction potential. As calculated by Gole, the A' and A" states are strongly repulsive at small Cl-OO angles. This is consistent with the large translational energy release that we observe in the Cl + O_2 products. It has already been noted that the 2A_2 \leftarrow 2B_1 transition does lead to considerable decrease in OClO bond angle. However, due to the increased bond length of the 2A_2 state, the expected O-O distance at the inner turning point for OClO 2A_2 bending is far too large to facilitate formation of an O=O bond. This suggests that the Cl + O_2 dissociation process must also involve strong coupling with one of the excited states of OClO. Two states are possible—the dark 2B_2

and the, as yet unobserved, ${}^{2}A_{1}$. Gole's calculations indicate equilibrium bond angles of $\sim 90^{\circ}$ for the ${}^{2}B_{2}$; that of the ${}^{2}A_{1}$ is near 120 degrees. Thus, coupling with the ${}^{2}B_{2}$ is also likely to play an important role in the Cl + O₂ dissociation mechanism.

We believe that more analysis of the present data combined the linewidth measurements will provide further insight into the photodissociation dynamics of OClO. We also hope that this work will inspire more detailed calculations on the ClO_2 and $Cl + O_2$ potential energy surfaces.

IV. CONCLUSIONS

We have clarified a number of questions regarding the formation of Cl + O_2 in the collision free photodissociation of OClO. Our results indicate that the yield is only appreciable (>1%) above 390nm. For a given energy, the yield of Cl + O_2 is always slightly greater under those lines corresponding to excitation of the symmetric bending motion, but is much smaller when exciting an asymmetric mode. The decrease in yield at shorter wavelengths or with asymmetric vibrational excitation likely results from faster predissociation to ClO + O. Suggested gas phase isomerization of OClO to ClOO, based on the earlier matrix isolation results of Arkell, *et al.*, ¹³ appears to be inappropriate. Although a substantial amount of O_2 is formed in the ground electronic state, there is strong evidence for a comparable yield for $O_2(^1\Delta)$, the total reaching a maximum of $3.9\pm0.8\%$ near 404nm.

According to Solomon, et al., 4,12 a 100% quantum yield for Cl + O₂ across the entire OClO absorption spectrum could lead to up to 30% of the observed Antarctic

ozone hole. Since we find that the quantum yield for $Cl + O_2$ in the collision free photodissociation of OCIO is not only small, but only extends over a relatively narrow portion of the absorption spectrum, we conclude that OCIO photodissociation in the gas phase plays a very minor role in the depletion of ozone currently seen in Antarctica. Further details regarding the mode specificity in the dynamics, complete yields of $Cl + O_2$ as a function of wavelength, and approximate ClO vibrational energy distributions will be presented in a forthcoming article.²⁴

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VII. FIGURE CAPTIONS:

- Fig.1: Energy level diagram for OCIO and possible photodissociation products.
- Fig.2: OClO absorption spectrum (204K), from Ref. 12. Vibrational assignments from Ref. 1.
- Fig.3: Newton diagram in velocity space for Cl-containing fragments from OClO photodissociation at 404nm. Arrow denotes initial OClO beam velocity, and circles are calculated recoil velocities for formation of ground vibrational state fragments.
- Fig.4: Product time-of-flight spectra (20°) recorded at m/e= 35 (Cl⁺) from indicated OClO vibrational levels. Inset shows expanded vertical scale (x15) to reveal structure in Cl + O₂ channel. Cl+O₂ yields are also indicated. ··· Experimental Data. -Optimized fit.
- Fig.5: Time-of-flight spectra (40°) for the O_2 and O atom channels recorded at m/e=32 (O_2^+) , and m/e= 16 (O^+) , from (5,1,0) level. Contribution to O^+ from fragmentation of O_2 is negligible and contribution from ClO is broad peak at $T=350-600\mu$ sec.
 - · · · Experimental Data. - Optimized fits based on Fig. 6.
- Fig. 6: Translational energy distributions for $Cl(^2P) + O_2$ and $O(^3P) + ClO$ channels from (5,1,0) level. The calculated maximum relative translational energies for production of internally excited diatomics are indicated.
- Fig.7: OCIO Absorption Spectrum, taken from Ref. 22. Also indicated are quantum yields for the $Cl + O_2$ channel from preliminary analysis of the data.

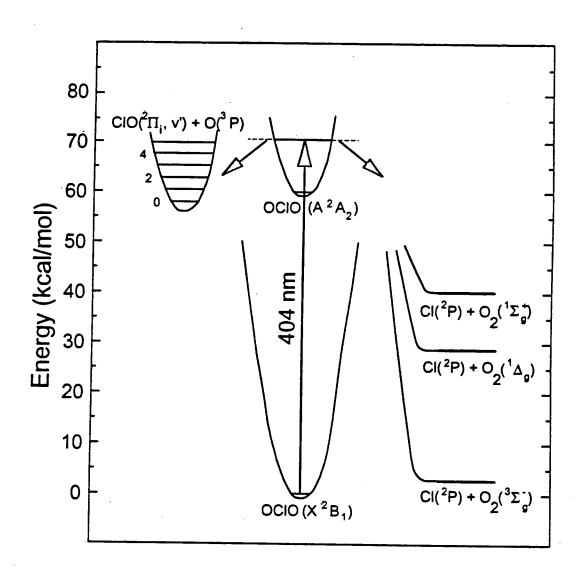


Fig. 1

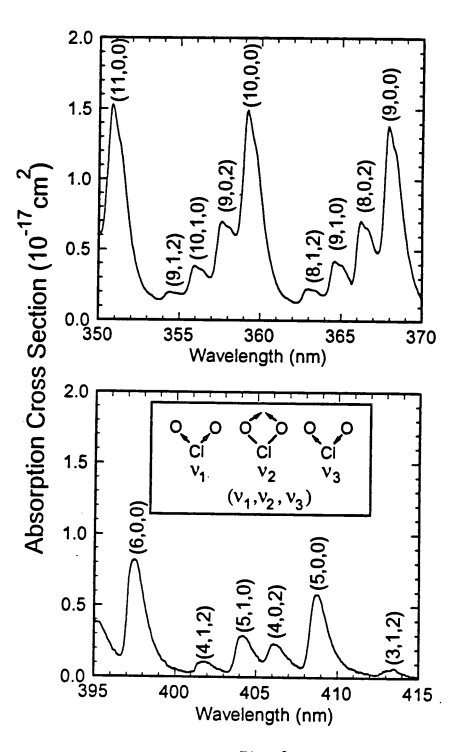


Fig. 2

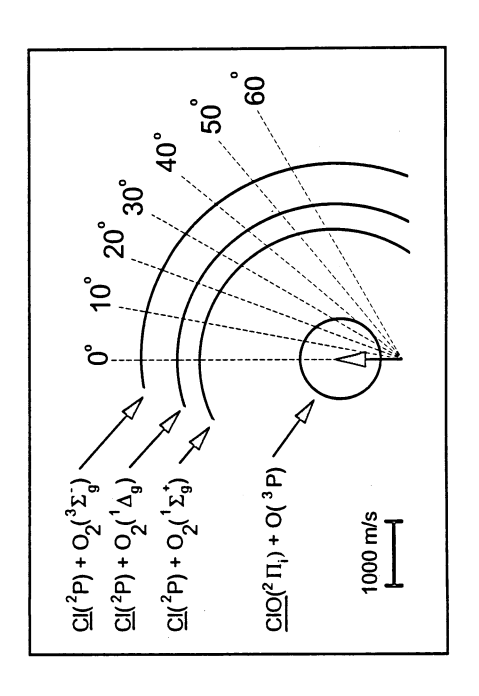


Fig. 3

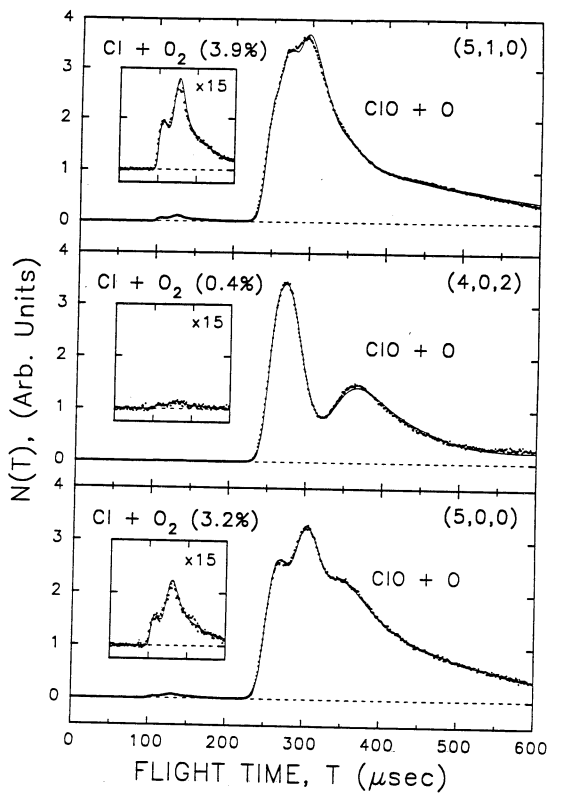


Fig. 4

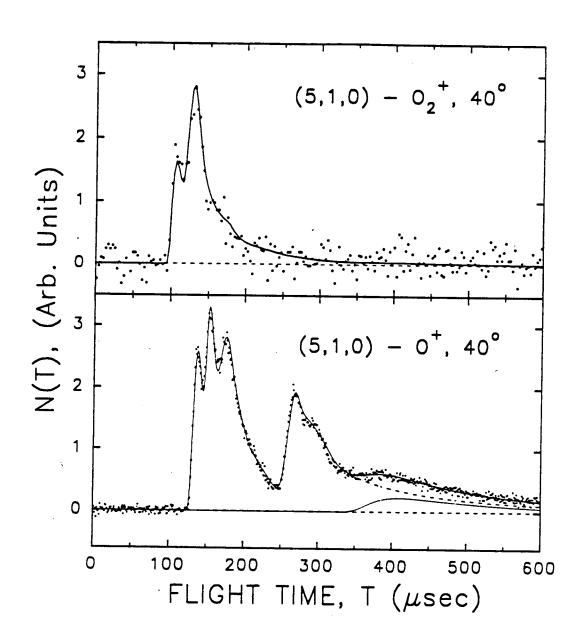


Fig. 5

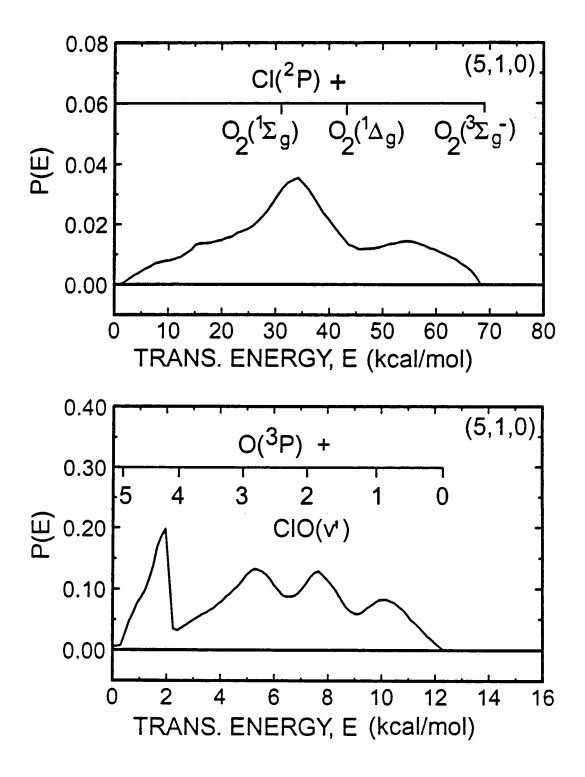
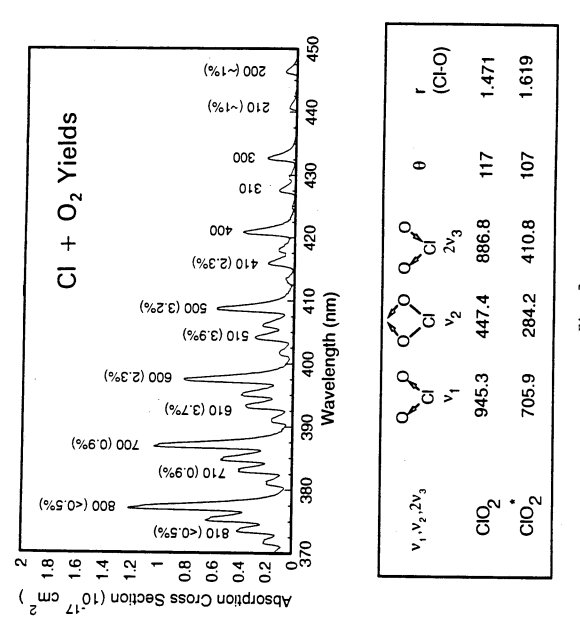


Fig. 6



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LAWRENCE BERKELEY LABORATORY
UNIVERSITY OF CALIFORNIA
TECHNICAL INFORMATION DEPARTMENT BERKELEY, CALIFORNIA 94720